HORMONE-DEPENDENT AND HORMONE-INDEPENDENT CONTROL OF METABOLIC AND DEVELOPMENTAL FUNCTIONS OF MALATE DEHYDROGENASE – REVIEW

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Malate dehydrogenases (MDHs) are specific class of ubiquitous and multimeric oxidative de-carboxylases with well conserved amino acid sequences in structurally important regions and with similar overall structural topology. They mostly use malate or oxaloacetate as substrates to generate pyruvate and utilize preferentially NADP or NAD as cofactor. Among species and even within an organism they differ in their subcellular localization and specificity for the cofactor. Comparison across microbial, plant and animal kingdoms show that MDHs were able to adopt tissue-, species- or environmental-specific functions while still keeping main structural features. Although basic principles of MDH regulation are similar to other enzymes and include oligomerization, cofactor binding, divalent cation availability, some of MDH enzymes are regulated also at different levels involving control of hysteresis, protein-protein interaction and gene expression. In this review we concentrate on those aspects of MDH function and regulation in animals that are specifically associated with cell differentiation and proliferation, ontogenic development, hormonal control, and partly with diseases. Accenting these aspects of MDHs provides emerging and new views on their regulatory function in complex eukaryotic metazoan organisms that goes beyond their classical role in basic metabolism.

Introduction

Malate dehydrogenases, or MDHs (EC 1.1.1.37/EC 1.1.1.38/EC 1.1.1.39), constitute a family of evolutionarily very old and ubiquitous multimeric enzymes that reversibly catalyze the conversion of oxaloacetate and malate into pyruvate, and that differ in their subcellular localization and specificity for the cofactor. The cytosolic MDHs (L-malate:NADP⁺ oxaloacetate-decarboxylating oxidoreductase; EC 1.1.1.40), also known as NADP-malic enzymes (ME), catalyze the NADP⁺ dependent oxidative decarboxylation of malate into pyruvate and carbon dioxide to generate NADPH. Due to its ability to produce NADPH, ME is thought to be a key enzyme in lipid biosynthesis. Comparison of phylogenetically distant MDHs reveals a relatively low degree of sequence similarity. However, several residues located in motifs and domains critical for substrate recognition, catalysis, cofactor binding and subunit interface are highly conserved (Tripathi 1993; Goward and Nicholls 1994; Goodridge et al., 1996; Hill et al., 1996; Escher and Widmer 1997; McAlester-Henn and Small 1997). Eukaryotic ME is found in both vertebrates and invertebrates as well as in plants. These enzymes require Mg²⁺ or Mn²⁺ for their activity and also display a high specificity and low Km for NADP⁺. At least for MDHs of the EC 1.1.1.37 family of enzymes it is known that they catalyze the conversion of oxaloacetate and malate, and this reaction is coupled with easily detectable cofactor oxidation/reduction. The MDHs are ubiquitous enzymes displaying several isoforms, for which several isoforms have
been identified, differing which differ in their subcellular localization and specificity for the cofactor NAD or NADP (GIETL 1992a).

Organelle-targeted malate dehydrogenases (mitochondrial, peroxisomal, chloroplast, and glyoxysomal) are synthesized with 25-55 amino terminal extension that is cleaved off in connection with the import of the enzyme precursor into the organelle (GIETL 1992b). Several of these organelle-specific malate dehydrogenases are remarkably similar with respect to their kinetic properties, but differ widely in other characteristics, e.g. molecular weight (MW), isoelectric point (pI), thermostability and serological properties (HOCK and GIETL 1982). Although not regular, but several MDHs display competitive affinity for two different cofactors. For example, some animal as well as plant NAD-malic enzymes were shown to possess intrinsic NAD-dependent malic enzyme activity. The NAD-dependent activity is competitively inhibited by micromolar concentrations of NADP and NADPH, reflecting the high affinity of malic enzyme for NAD(P)H. Several observations on these enzymes indicate that the NAD- and NADP-dependent activities are likely catalyzed by the same active site. Manipulating divalent cations and the use of NAD as an alternating coenzyme have revealed interaction between the binding of coenzyme and metal ions and their effects on the Km values of each of the other participants in the malic enzyme reaction. The affinity of malic enzyme for the divalent metal ions like Mg\(^{2+}\) and particularly Mn\(^{2+}\) or even for alternative substrate was also shown do be dependent on the nucleotide coenzyme. In turn, the divalent metal ion influences the affinity of the enzyme for the coenzyme as well as substrate (ASHTON 1997).

In general our present knowledge about MDH proteins reflects past interest in their role in basic metabolism with special emphasis on cytosolic-mitochondrial metabolite shuffling, indirect control of tricarboxylic acid (TCA) cycle, involvement in glycolytic and oxidative pathways, and lipogenesis. These aspects are discussed in detail in several reviews (GIETL 1992a; TRIPATHI 1993; GOWARD and NICHOLLS 1994; HILL et al., 1996; MCAULISTER-HENN and SMALL 1997; MUSRA\v{T} et al., 1998; MEVARE\v{C} et al., 2000; DRINCOVI\v{C} et al., 2001; MARIIUENDA-EGEA and BONETE 2002; MINARIK et al., 2002; TEHEI and ZACC\'AI 2005) which we recommend to the attention of interested readers. In spite of the fact that some of the MDHs are able to respond to imbalanced physiological conditions by changes in enzyme activity and could also serve as potential differential diagnostic tools, surprisingly there is limited knowledge on misfunctioning MDHs in diseases or adverse physiological conditions. One of the reasons why it is so may be extreme sensitivity of the organism to altered (mutated) or simply misfunctional MDH and, therefore, most of such a mutational changes in MDH enzymes were eliminated by selection pressure due to their lethal consequences. Based on some recent data and results that have not been in the centre of the views, here we would like to accent several aspects of MDH function that go beyond the basic metabolism and deal with their potential role in cell differentiation and ontogenic development thus providing emerging and new views on their regulatory function in eukaryotic metazoan organisms.

**Differentiations state-specific and disease-related functions of MDHs**

There are several indications that malate dehydrogenases may reflect growth phase or differentiation state of a particular tissue. For example, insulin-induced early growth phase in hepatic cells is accompanied by short-term repression of cytosolic ME (BARROSO and SANTISTEBAN 1999). Growth phase and differentiation state of cells is particularly well studied in malignancies and transformed cells. It has been found that poorly differentiated malignant cells show high aerobic glycolysis. Also other metabolic ways are deficient, as for example, intramitochondrial aldehyde catabolism, at the origin of a possible acetaldehyde toxicity, which can be circumvented by the synthesis of an unusual and neutral product for mammalian cells acetoin, through tumoral pyruvate dehydrogenase. If most of the glycolytic pyruvate is deviated to lactate production, little of the remaining carbons enter a truncated Krebs cycle where it feeds sterol synthesis. Glutamine is the major oxidizable substrate by many tumor cells. Inside the mitochondrion, it is deaminated to glutamate through a phosphate-dependent glutaminase (BAGGETTO 1992). Under these conditions, glutamate is preferentially transaminated to \(\alpha\)-ketoglutarate that enters the Krebs cycle. Glutamine may be completely oxidized through the abnormal Krebs cycle only if a way of forming acetyl CoA is present: cytosolic malate entering mitochondria is preferentially oxidized to pyruvate and \(\text{CO}_2\) through an intramitochondrial NAD(P)\(^{+}\)-malic enzyme (ME), whereas intramitochondrial malate is preferentially oxidized to oxaloacetate through malate dehydro-
genase, thus providing a high level of intramitochondrial substrate compartmentation (Baggetto 1992). These observations indicated that described and perhaps many other regulatory aberrations in tumor cells appear to be reflections of a complex set of non-random phenotypic changes, caused by misexpression of oncogenes and/or oncogene-driven genes. Although not a primary cause, several of these metabolic aberrations could serve as diagnostic marker in some types of cancer and other diseases.

When analyzing mitochondrial and cytosolic NADP⁺- and NAD⁺-dependent malic enzymes in several human tumor cell lines it became evident that both mitochondrial and cytosolic NADP⁺-ME have significantly increased activity (Balinsky et al., 1983; 1984; Loebber et al., 1994). On the other hand, earlier studies suggested that the deficiency of cytosolic ME is a possible marker for malignancy in lymphoid cells (Pover et al., 1981). This conclusion cannot be extended to tumor cells derived from tissues of epithelial origin, since all carcinoma cell lines studied so far have considerable NADP⁺-ME activities. An extended study investigating the status of the different MEs in a large number of human cell lines has shown that no mitochondrial NADP⁺-ME was found in the tumor cell lines tested. This finding differs from a report of an increase in mitochondrial NADP⁺-ME of preneoplastic and tumor-derived rat tracheal cells when compared to corresponding normal primary cells (Wasilenko and Marchok 1985). However, it is possible that this reported increase in NADP⁺-ME activity reflected mitochondrial NAD⁺-ME since, in contrast to the human NAD⁺-ME which is hardly active with NADP⁺ (Sauer and Dauchy 1978; Sauer et al., 1980), rat NAD⁺-ME has significant activity when assayed with NAD⁺ (Sauer and Dauchy 1978; Loebber et al., 1991). By contrast, breast cancer tumor cells show reduced activity in mitochondrial ME (Perumal et al., 2005) which was often characterized as pI 7.8 isoform (Mazurek et al., 1996, 1998, Caetano et al., 1997).

Several studies have proved that the cytosolic NADP⁺-ME expressed in the tumor cells is the same as the ME expressed in normal liver and adipose tissue (Sauer and Dauchy 1978; Loebber et al., 1991). Posttranslational modifications specific for tumor cells affecting the activity of the protein are unlikely since the recombinant ME has the same size, as seen in SDS-polyacrylamide gels. In normal tissues, cytosolic ME provides NADPH for fatty acid biosynthesis. However, it has been found that the NADPH production by ME was small when compared to glucose-6-phosphate dehydrogenase or NADP⁺-dependent isocitrate dehydrogenase the activity of which is about 50% cytosolic (Sauer and Dauchy 1978; Loebber et al., 1991; Noguchi et al., 1993). It is unlikely that cytosolic ME contributes significantly to the NADPH pool in the cytoplasm of the tumor cells. It is more likely that pyruvate is the relevant product of the ME reaction in tumor cells.

In transformed oval liver epithelial cells that proliferate during the early stages of hepatocarcinogenesis, a strong correlation between the glycolytic flux rate and glutamine consumption as well as glutamate production was observed. Furthermore, the transport of glycolytic hydrogen, produced by the glyceraldehyde 3-phosphate dehydrogenase-catalyzed reaction, from the cytosol into the mitochondria by means of the malate-aspartate shuttle was enhanced, due to alterations in the activities of malate dehydrogenase and glutamate oxaloacetate transaminase. The upregulation of the glycolytic hydrogen transport and the alterations in the glycolytic enzyme complex led to an enhanced pyruvate production at high glycolytic flux rates, thus affecting cytosolic malate dehydrogenase activity at the level of end product (Mazurek et al., 1999). This indicated that a special metabolic feature, in this case increased glycolysis and glutaminolysis, is characteristic for tumor cells.

Human cytosolic malate dehydrogenase (so-called MDH1) is important in transporting NADH equivalents across the mitochondrial membrane, controlling tricarboxylic acid (TCA) cycle pool size and providing contractile function. Cellular localization studies indicate that MDH1 mRNA expression has a strong tissue-specific distribution, being expressed primarily in cardiac and skeletal muscle and in the brain, at intermediate levels in the spleen, kidney, intestine, liver, and testes and at low levels in lung and bone marrow (Lo et al., 2005). The variety of MDH functions are primarily related to aerobic energy production for muscle contraction, neuronal signal transmission, absorption/resorption functions, collagen-supporting functions, phagocytosis of dead cells, and processes related to gas exchange and cell division. During neonatal development, MDH1 is expressed in human embryonic heart as early as the 3rd month and then become upregulated from the 5th month until the birth. The expression of MDH1 is maintained in the adult heart but is not present in levels as high as in the fetus. Interestingly, highly overexpressed levels of MDH1 are found in left ventricular cardiac muscle of dilated cardiomyopathy.
(DCM) patients when contrasted to the diseased non-DCM and normal heart muscle by in situ hybridization and Western blot. These observations indicate that changes in MDH1 activity or expression appears to be related to activation of glucose oxidation in relatively hypoxic environments of fetal and hypertrophied myocardium (Lo et al., 2005). Very recent studies of Yoon et al. (2006) demonstrated that mammalian cytosolic malate dehydrogenase named Mor2 plays even more crucial role in mice oocyte maturation and embryo development. Silencing Mor2 function by RNA interference in germinal vesicle- and metaphase I (MI)-arrested oocytes has greatly decreased oocyte maturation as well as the rate of blastocyst development. Thus, cytosolic MDH is an essential factor for oocyte maturation and embryo development in mouse most probably by providing NADH and controlling TCA cycle (Yoon et al., 2006).

The significance of mitochondrial NAD+-ME is thought to be related to the use of glutamine as a respiratory fuel, in a pathway from glutamine to lactate that has been named glutaminolysis (McKeehan 1982). This pathway not only provides energy by using a truncated Krebs cycle from α-ketoglutarate to malate, it also produces intermediates for anabolic purposes in tumor cells, e.g. aspartate and nitrogen groups for DNA biosynthesis. Malate generated from glutamine in the mitochondria has three possible options: conversion to oxaloacetate by malate dehydrogenase, decarboxylation to pyruvate in the mitochondria by NAD+-ME, or transport into the cytoplasm. Here, malate can be decarboxylated by cytosolic ME with subsequent conversion to lactate by lactate dehydrogenase. NAD+ produced by this reaction can be used by glyceraldehyde-3-phosphate dehydrogenase, thus compensating for the expression of a partially inactive pyruvate kinase in tumor cells (Eigenbrodt et al., 1992). A significant proportion of glutamine taken up by tumor cells and lymphocytes ends up in lactate, which is secreted (McKeehan 1982; Eigenbrodt et al., 1985; Brand et al., 1989). Therefore, a possible function of the cytosolic NADP+-ME in tumor cells is the conversion of excess mitochondrial malate, which can cross the mitochondrial membrane readily, to pyruvate, which is subsequently converted to lactate and extruded from the tumor cell.

It is becoming well established (more evident) that alterations in the activities of enzymes related to energy metabolism (Washizu et al., 2005) in malignant cells or cells undergoing the process of transformation are due to alterations in the regulation of mitochondrial and cytosolic MDHs, cytosolic lactate dehydrogenase (LDH) and often also pyruvate kinase (PK). In many cases this process leading to changes in energy metabolism is started by misregulation of one or few genes coding for enzymes participating in TCA or closely related metabolic cycle (Rao et al., 1997; Arai et al., 2002a,b). Subsequently, misbalanced substrate/product ratio alters activity of other enzymes that finally can be observed as changes in the activity of several other enzymes as well as changes in their mutual ratio (aerobic oxidative versus anaerobic metabolism). Frequently glycolytic rate increases, more pyruvate goes into the Krebs cycle than into lactate. The availability of glucose-derived pyruvate for oxidative metabolism would mean less of a dependency on glutamine as a carbon source (Ross and Medina 1998; Ross et al., 2000). Clearly, alterations in mitochondrial versus cytosolic enzyme activities and their ratio expressed as rate between aerobic oxidative and anaerobic metabolism can now be considered to be not only a good indicator of energy metabolism related to glucose utilization and aid in understanding of tumor metabolism but also with expression patterns of selected markers can also serve as potentially excellent diagnostic tools for the status, response to chemotherapy and prognosis of malignancy.

Hormonal regulation of MDH functions

Numerous studies in mammals and other vertebrates show that MDH activity can be co-operatively controlled by nutrients and hormones. Vertebrate MDH is a target for action of several hormones: catecholamines, insulin, thyroid hormone and retinoic acid (Ballard and Hanson 1967; Geer et al., 1976, 1980; Towl et al., 1980; Dozin et al., 1985; Knopp and Brøtø 1987; Morioka et al., 1989; Hernandez et al., 1991; Mogensen et al., 2007). It is well known that in mammalian and avian models, thyroid hormone (Ortiz-Caro and Jolin 1991; Mooradian et al., 1991; Sali et al., 1991; Mann et al., 1992) and some steroids like estradiol (Astiazaran et al., 1989; Patnaik, 1990), androsterone (Song et al., 1989; Mohan and Cleary, 1991) and nandrolone (Tyllicki et al., 2007) may stimulate activity of MDH by increasing MDH gene transcription and stabilizing MDH mRNA (Song et al., 1988; Desvergne et al., 1991). Thyroidal control of malate dehydrogenase, specifically cytosolic malic enzyme, activity and function is well documented phenomenon in higher vertebrates (Frenkel 1975; Goodridge 1975; Bagchi et al.,...
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1986; Goodridge et al., 1986, 1989, 1991, 1996; Nikodem et al., 1989; Iritani 1992). Moreover, rat MDH is positively regulated by both thyroid hormones and retinoids via their nuclear receptors recognizing specific response elements upstream of MDH promoter (Morioka et al., 1989; Petty et al., 1990). In this case hormonal regulation crosstalks with factors of nutritional pathway to control lipogenesis at the level of malic enzyme that generates NADPH. Besides MDH and especially ME also additional enzymes such as glucose-6-phosphate dehydrogenase (G6PDH) and 6-phosphogluconate dehydrogenase (6PGDH) can be upregulated by thyroid hormones (Varghese et al., 2001).

In chick embryo hepatocytes in culture, insulin and triiodothyronine (T3) were identified as positive effectors and glucagon, acting via cyclic AMP (cAMP), as a negative effector of ME activity. Hormone concentrations in blood are consistent with insulin and T3 playing the major positive roles, and glucagon a major negative endocrine role, in regulating hepatic malic enzyme activity during the transitions between the fed and the starved states. It was found that not only insulin but insulin-like growth factor 1 also stimulates accumulation of malic enzyme. Nutrition- and hormone-induced changes in malic enzyme activity are due to altered concentrations of ME protein which, in turn, are due to altered rates of synthesis of ME. Synthesis of ME is controlled by regulating the level of ME mRNA which, in turn is regulated at initiation of transcription (Goodridge et al., 1989).

Further experimental evidence obtained in mammals show that a low concentration of cAMP can act as a second messenger in a crosstalk between steroid hormones and retinoid acid resulting in the stimulation of a second messenger in a crosstalk between steroid hormones and retinoid acid resulting in the stimulation of cAMP is known to be involved in mediating hysteretic behavior of ME which was described for many oxidoreductases including MDH and ME (Soulie et al., 1988; Aon et al., 1989; Cebon et al., 1990; Edwards and Andreo 1992; Avilan and Garcia, 1994). Mobilization of existing pools of the enzyme can be achieved by several mechanisms including posttranslational modification or releasing enzyme protein from inactive state e.g. from interaction with chaperons etc.

Responsiveness of MDH or ME to hormonal stimuli is not always necessarily associated with lipogenesis, but rather it can reflect intensity of actual and sustaining demands of tissue specific energy metabolism. Thus, given the central role of the heart in responding to physiological changes and the high energy requirements of the myocardium, it is not surprising that cardiac myocytes are sensitive to the effects of thyroid hormone (Ralphie et al., 2004). Thyroid hormone acts on the heart through mechanisms that are both independent of and dependent on gene transcription. Acute effects seen early after exposure to the bioactive form of thyroid hormone, T3, include increases in heart rate, ejection fraction, and cardiac output, as well as increases in blood volume (Klein 1988). Important myocyte genes responsive to thyroid hormone that impact mechanical function have been identified, including α-myosin heavy chain, sarcoplasmic reticulum Ca2+-ATPase, Na+-K+-ATPase, and several voltage-gated potassium channels (Pantos et al., 2004). Beyond its effects on cardiac mechanical function, thyroid hormone also directly impacts cardiac energetics and mitochondrial function. Thyroid hormone actions on mitochondria result from direct regulation of both nuclear and mitochondrial DNA transcription as well as from hemodynamic changes that indirectly alter gene expression (Gomberg-Maitland and Frishman, 1998). Several nuclear-encoded respiratory chain genes have been identified that are induced after administration of T3, including cytochrome c1, cytochrome-c oxidase, F1-ATP synthase subunit, glycerol-3-phosphate dehydrogenase, and MDH (Goglia et al., 1999). In addition, the nuclear action of T3 on gene transcription, and energy metabolism can be further enhanced by facilitating mitochondrial protein import pathway including mitochondrial MDH (Colavecchia et al., 2003).

Control of energy metabolism by regulating mitochondrial
MDH activity was found also in atypical situations associated with idiopathic pulmonary arterial hypertension (IPAH), a pathology related to low levels of the vasodilator nitric oxide (NO), that appears to be coregulated by insulin, thyroid and steroid hormones (Xu et al., 2007).

Molecular mechanisms controlling hormonal response of MDH/ME genes are of central interest for several years and it is still current issue as not all crucial questions were successfully answered. For example, in liver insulin stimulates the transcription of the gene encoding the cytosolic form of ME and modulates protein binding to two putative insulin response sequences (IRSs) in the ME promoter. One of these IRSs resembles that identified in the phosphoenolpyruvate carboxykinase (PEPCK) gene, whereas the other resembles that defined in the glyceraldehyde-3-phosphate dehydrogenase (GAPDH) gene. To assess the functional significance of these changes in protein binding, a series of truncated ME-chloramphenicol acetyltransferase (CAT) fusion genes were transiently transfected into rat H4IIE hepatoma cells. Deletion of the PEPCK-like IRS motif had no effect on the stimulation of CAT expression by insulin. Instead, the stimulatory effect of insulin was mediated through an AP-1 motif and an Egr-1 binding site that overlaps the GAPDH-like IRS motif. Both the ME AP-1 motif and the AP-1 motif identified in the collagenase-1 gene promoter were able to confer a stimulatory effect of insulin on the expression of a heterologous fusion gene, but surprisingly only the latter was able to confer a stimulatory effect of phorbol esters. Instead, the data suggest that AP-1 binds the ME AP-1 motif in an activated state such that phorbol ester treatment has no additional effect. The collagenase and ME AP-1 motifs were both shown to bind mainly Jun D and Fra-2, with similar affinities (Streeper et al., 1998). In chick embryo hepatocytes, activation of ME gene transcription by T3 is mediated by a T3 response unit (T3RU) that contains five T3 response elements (T3REs) plus five accessory elements that enhance T3 responsiveness conferred by the T3REs. Results from in vitro binding assays indicate that one of the accessory elements (region F) binds CCAAT/enhancer-binding protein-alpha (C/EBPα). Transfection analyses demonstrated that the stimulation of T3RE function by region F did not require the presence of additional malic enzyme gene promoter sequences. Expression of a dominant negative C/EBP inhibited the ability of region F to stimulate T3 responsiveness. In chromatin immunoprecipitation assays, C/EBPalpha and TR associated with the malic enzyme T3RU in the absence and presence of T3 with the extent of the association being greater in the presence of T3. These observations indicate that C/EBPalpha interacts with TR on the malic enzyme T3RU to enhance T3 regulation of ME gene transcription. T3 treatment increased the acetylation of histones, decreased the recruitment of nuclear receptor corepressor and increased the recruitment of steroid receptor coactivator-1, CREB binding protein, and the thyroid hormone associated protein/mediator complex at the malic enzyme T3RU. In contrast, T3 treatment had no effect on the acetylation of histones and the recruitment of corepressors and coactivators at the T3RU that mediates the T3 activation of acetyl-CoA carboxylase-α gene transcription (Yin et al., 2005). These observations indicate that ME activation by T3 can be, at least in part, C/EBPα-dependent, although there are several other pathways that converge onto C/EBPα protein which is becoming an important crosstalk point. There is an evidence, that T3 transcription of ME in hepatocytes can be further enhanced by cooperative binding of chicken ovalbumin upstream-promoter transcription factor (COUP) and E-box-binding proteins to own enhancer within ME promoter (Wang et al., 2002). Recently a novel function has been assigned to Spot14 (S14) protein the expression of which has been found to be co-responsive with ME to T3 long time ago but leaving its biochemical function elusive (Goodridge et al., 1987). Experiments of Chou et al. (2007) have shown that this small acidic protein forming homodimer is a nuclear factor that functionally interacts with thyroid hormone receptor (TR), a mediator of T3 action and can act as a positive or negative cofactor of the TR to regulate ME gene expression.

### Developmental aspects of MDH action and functions

Although lipogenesis-prone or independent enzyme activity is continuous process during the life span of many eukaryotes, its intensity often displays developmental, cell cycle or ageing dependent variations. To study such developmentally-linked variations several model organisms have been used in which the activity of lipogenic enzymes including MDHs has been analyzed at biochemical or molecular level (Ballard and Hanson 1967; Geer et al., 1976; Towle et al., 1980; Morioka et al. 1989). This indicates that regardless of lipogenesis MDH enzymes not only reflect ongoing changes in ontogenic phases but may play a specific role in ontogenic development.
The procyclic forms of *Trypanosoma brucei* parasite possess three different malate dehydrogenases (mitochondrial, glycosomal and cytosolic). The latter is the only malate dehydrogenase expressed in the bloodstream forms, thus confirming that the expression of malate dehydrogenase isozymes is regulated during the *T. brucei* life cycle. Cloning followed by their expression in Escherichia coli cultures has revealed that mitochondrial malate dehydrogenase showed to be more active than glycosomal malate dehydrogenase in the reduction of oxaloacetate. Nearly 80% of the total activity in procyclic crude extracts corresponds to the former isozyme which also catalyzes, although less efficiently, the reduction of p-hydroxyphenyl-pyruvate. Importantly these studies helped to distinguish *Trypanosoma brucei* from *Trypanosoma cruzi*, since in the latter parasite a cytosolic malate dehydrogenase is not present and mitochondrial malate dehydrogenase specifically reduces oxaloacetate which can be very useful in diagnosis (Aranda et al., 2006).

Quite surprisingly gene encoding malic enzyme (ME) in water arthropod, copepod *Tigriopus californicus*, appears to play role in separation of its two divergent populations (Willett and Berkowitz 2007). Comparison of cloned genes has shown that each ME gene has diverged extensively between *T. californicus* populations and one gene shows evidence for a recent selective sweep. Segregation patterns of genotypes for both ME genes in adult F2 hybrids reveal dramatic departures from Mendelian inheritance. Some developmentally programmed changes in NADPH-associated enzymes of rat liver including MDH, succinate dehydrogenase, glutamate dehydrogenase, glucose-6-phosphate dehydrogenase can be reprogrammed during embryogenesis. In utero exposure to hydroxyprogesterone leads to significantly increased levels of these enzymes in adult male livers, while lactate dehydrogenase (LDH) activity was greatly increased. Simultaneously, while lactate dehydrogenase (LDH) activity was greatly decreased in newborn myocardium. These observations show that myocardial malate-aspartate shuttle capacity declines rapidly after birth, and indicate that divergence of mitochondrial MDH mRNA and protein levels during development could be due to posttranscriptional regulation of this protein (Scholz et al., 1998). Results also suggest that developmental decline in malate-aspartate shuttle capacity appears to be regulated by decreased oxoglutarate/malate carrier gene expression. The malate/aspartate as well as α-glycerophosphate shuttles show existence of sufficient capacity in cardiac mitochondria to accommodate increased shuttle flux even in hypertrophied myocardium that becomes more glycolytically active (Rupert et al., 2000). However, elevated thyroid hormone leads to upregulation of metabolic pathways associated with energy production and development of heart hypertrophy, as observed by Ralph et al. (2005). Expression of excitatory amino acid transporter type 1 (EAAT1)
mRNA and protein that functions as a glutamate carrier in the malate/aspartate shuttle was increased nearly threefold in T3-treated animals, whereas expression of aralar1 and citrin (both cardiac mitochondrial aspartate-glutamate transporters) mRNA and protein levels remain decreased and unchanged, respectively. On the other hand, hypothyroidism resulted in a decrease in EAAT1 mRNA, but without any effect on the EAAT1 protein level or shuttle activity. This partially hormonal regulation of EAAT1 may facilitate increased metabolic demands of the cardiomyocyte during hyperthyroidism and impact cardiac function during development of hyperthyroidism.

Early neonatal and adult development of rat brain is accompanied by ontogenic changes in the cerebral pyruvate recycling pathway and the cellular localization of associated enzymes, ME and phosphoenolpyruvate carboxykinase (PEPCK). In contrast to gradually decreasing level of PEPCK, ME activity is steadily increasing to adult levels. Both enzymes are expressed in astrocytes and in synaptosomes, but cortical neurons showed only PEPCK activity but no detectable ME activity (Cruz et al., 1998). For developmentally-linked changes of human placental MDH activity is typical their variations related to levels of palmitoyl-CoA and oleate. As was shown by Bandypadhyay et al. (1994) these changes can rather reflect modulating activity of fatty acid binding proteins (FABPs), an important mediator of fatty acid transport and action. Its function is rather peculiar as FABPs enhance the activity of malate dehydrogenase in the presence of palmitoyl-CoA or oleate and also protect against palmitoyl-CoA or oleate inhibition. Ontogenic related MDH activity is detected throughout the gestation, showing a peak at midgestation, and it is anticipated that the modulatory effect of FABP may be due to the binding of long chain fatty acyl-CoA or fatty acid rather than a direct effect of FABPs on the enzyme.

Also development and capacitation of mammalian sperm are accompanied by expression of several marker proteins including MDH and isocitrate dehydrogenase (representatives of TCA cycle), and several other proteins like lactadherin P47, acrosomal protein SP-10 precursor, prohibitin, DJ-1 protein, arylsulfatase-A, and cytochrome c (Choi et al., 2008). At least expression of cytochrome c appears to be regulated via p53 pathway and may be involved in decision developmentally-linked programmed cell death or further spermiogenesis. If developmental decision is switched towards apoptosis, MDH and isocitrate dehydrogenase are rapidly downregulated.

Besides above mentioned cases, there are additional non-lipogenic functions or unconventional roles of MDHs. Skeletal muscle development is accompanied by steadily but modestly increased levels of mitochondrial MDH, ATP synthase, succinate dehydrogenase, isocitrate dehydrogenase, pyruvate kinase, enolase and phosphofructokinase. Unexpectedly, in the process of ageing and under some pathological situations like sarcopenia, senescent muscular fibres show significantly increased levels of MDH, succinate dehydrogenase, MDH, ATP synthase, succinate dehydrogenase, isocitrate dehydrogenase and isocitrate dehydrogenase, whereas pyruvate kinase, enolase and phosphofructokinase are apparently downregulated (Doran et al., 2008).

Some examples of MDH regulation and involvement in various processes described above may appear as episodic but gradually accumulating evidence from other model organisms, e.g. insects including Drosophila, provides more support for developmentally-linked role of malate dehydrogenases. In Drosophila melanogaster, several oxidative NADP-enzymes including cytosolic MDH (ME) have been extensively studied. It has been shown that activity of ME is low at the beginning of postembryonic development, gradually increases during larval life peaking in the first half of the last larval instar, is low during the pupal stage and freshly eclosed adults, then rises again with increasing age of adults (Redkin 1970; Chernik et al., 1982; Kaplin and Korochkin, 1987; Geer et al., 1978a). The course of ME activity during larval development shows a strict pattern but it may be modulated by dietary factors. High saccharide diet results in increase of ME activity in feeding larvae while low saccharide/high lipid diet makes ME activity drop to almost undetectable levels (Geer et al., 1976, 1978b, 1980; Geer and Perille 1977). Also, there is a tissue specific distribution of ME in Drosophila body with highest activity in larval fat body and a significant portion in larval imaginal discs (Kuhn and Sprey 1987) and adult oviduct, ejaculatory duct and paragonia (Pinkröhner et al., 1985). Later it was found that Drosophila ME is regulated by juvenile hormone (JH) and JH analogues. However, the response of ME to JH depends on the developmental stage of the animal in relation to its endogenous titre of ecdysteroid hormones. During the interecdysial period of the last instar, when endogenous ecdysteroids are low, ME rapidly responded to JH by increasing activity, while little or no response was monitored in wandering (post-feeding) larvae, i.e., after a pulse of ecdysteroids. Activity of ME in ecdys-
teroid-deficient mutants of *Drosophila* increased during a shift to restrictive temperature as compared to wild-type controls, and was sensitive to administration of JH. In contrast, when ecdysteroid-deficient mutants larvae during restrictive conditions were fed on a 20-hydroxyecdysone diet, JH was unable to increase the activity of the ME enzyme. When JH was applied prior to 20-hydroxyecdysone feeding, the activity of ME was significantly stimulated. Two phases of ME response to JH, dependent on ecdysteroids, could be observed. The first phase was short and independent of RNA and protein syntheses. But activation of ME in later stages of the response was inhibited by actinomycin D and cycloheximide, indicating that ME might be regulated by JH at the transcriptional and/or translational level (FARKAŠ and KNOPP, 1997). Another *Drosophila* mutant, *ap*4 which is naturally deficient in JH production, displayed significantly reduced activity of ME in heterozygotic combination, and almost undetectable ME activity in null homozygote adults. The *ap*4 phenotype was more strongly manifested in the adult stage than in larvae which showed 7-times lower requirements for JH titre. In addition, high/low sucrose diet fed to wild types or mutants affected the activity of larval ME, but the enzyme remained sensitive to administration of JH. These results corroborate those described for mammals and provided the first evidence that *Drosophila* ME might be under differential hormonal and nutritional control (FARKAŠ and KNOPP, 1998).

Further studies, also involving *ecd*1 and *su(f)*ts67g ecdysteroid-deficient mutations, revealed that response of ME to JH requires the presence of a minimal level of the steroid hormone ecdysone, showing a complex hormonal regulatory circuit in the execution of the JH response (FARKAŠ et al., 2002). Recently positive responsiveness of orthologous ME in a silkworm, *Bombyx mori*, to vertebrate steroid estradiol-17beta (E2) has been reported. The E2 has been described as one of vertebrate steroid hormones that have been identified in invertebrates, but without known function. E2-induced increase in fat body enzyme activity in 5th instar larval females could be counteracted by simultaneous application of specific E2-inhibitor, supporting possibility that E2 can have an important metabolic function (ROY et al., 2007). Molecular mechanisms of many of these responses mentioned above share common principles and therefore we can see clear links between their vertebrate and invertebrate counterparts. By using appropriate molecular-genetic tools and model organisms hopefully will not remain elusive for a long time.

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