ROLE OF ISOVALERYL-CoA DEHYDROGENASE AND
SHORT BRANCHED-CHAIN ACYL-CoA DEHYDROGENASE
IN THE METABOLISM OF VALPROIC ACID: IMPLICATIONS FOR
THE BRANCHED-CHAIN AMINO ACID OXIDATION PATHWAY

Paula B.M. Luís, Jos P.N. Ruiter, Lodewijk IJlst,
Isabel Tavares de Almeida, Marinus Duran, Al-Walid Mohsen, Jerry Vockley,
Ronald J.A. Wanders and Margarida F.B. Silva

Research Institute for Medicines and Pharmaceutical Sciences – iMED.UL, Faculty of Pharmacy, University of Lisbon, Portugal (P.B.M.L.; I.T.A.; M.F.B.S.); Department of Clinical Chemistry and Pediatrics, AMC, Amsterdam, Netherlands (P.B.M.L.; J.P.N.R.; L.I.; M.D.; R.J.A.W.); Department of Pediatrics, School of Medicine, University of Pittsburgh, USA (A-W.M.; J.V.).

**Running title:** Valproate and the metabolism of branched-chain amino acids.

**Corresponding author:** 

Margarida Maria Fernandes Baptista e Silva

Research Institute for Medicines and Pharmaceutical Sciences -iMED.UL,

Faculdade de Farmácia da Universidade de Lisboa,

Av. Prof. Gama Pinto, 1649-003 Lisboa, Portugal

Tel: 351-217946400, ext 14511; Fax: 351-217946491; E-mail: mbsilva@ff.ul.pt

Number of text pages: 14;

Number of tables: 1;

Number of figures: 6;

Number of references: 30;

Number of words in the abstract: 249;

Number of words in the introduction: 580;

Number of words in the discussion: 762

Abbreviations: VPA: 2-n-propylpentanoic a cid o r va lproic a cid;  $\Delta^{2(E)}$ -VPA: 2-n-propyl-2-

pentenoic acid; Co A: co enzyme A; DephCoA: dephosphocoenzyme A; BCAAs: b ranched-

chain amino acids; VP-CoA: valproyl-CoA; VP-DephCoA: valproyl-dephosphoCoA.

Abbreviations of enzymes: IVD: Isovaleryl-CoA de hydrogenase (EC 1. 3.99.10); I BD:

Isobutyryl-CoA dehydrogenase (EC 1.3.99.3) and SBCAD: Short branched-chain acyl-CoA

dehydrogenase (EC 1.3.99.12).

### **ABSTRACT**

Many b iological systems including the oxi dative catabolic pathway for b ranched-chain amino ac ids (B CAAs) are af fected in vivo by valproate t herapy. In t his s tudy we investigated the potential effect of valproic acid (VPA) and some of its metabolites on the metabolism o f BCAAs. In vitro studies w ere pe rformed us ing isovaleryl-CoA dehydrogenase (IVD), iso butyryl-CoA de hydrogenase (IBD) a nd sho rt br anched-chain acyl-CoA dehydrogenase (SBCAD), e nzymes i nvolved in the d egradation pa thway o f leucine, va line and isoleucine. The enzymatic activities of the three purified hu man enzymes we re m easured us ing optimized HPLC p rocedures and the r espective ki netic parameters were determined in the absence and presence of VPA and the corresponding CoA and dephosphoCoA conjugates. Valproyl-CoA and valproyl-dephosphoCoA inhibited IVD activity significantly by a purely competitive mechanism with  $K_i$  values of 74±4  $\mu$ M and 17 0±12 µM, respectively. IBD activity was not affected by any of the tested VPA esters. H owever, v alproyl-CoA did in hibit S BCAD activity by a pur ely competitive mechanism w ith a K<sub>i</sub> of 249±29 μM. I n addition, va lproyl-dephosphoCoA inhibited SBCAD activity via a distinct mechanism ( $K=511\pm96 \mu M$ ) which appeared to be of the mixed type. Furthermore, we also show that both SBCAD and IVD are active using valproyl-CoA as a substrate. The catalytic efficiency of SBCAD turned out to be much higher than of IVD, demonstrating that SBCAD is the most probable candidate for the first dehydrogenation step of V PA β-oxidation. Our da ta e xplain some of t he e ffects of valproate on the branched-chain a mino acid metabolism and shed new light on the biotransformation pathway of valproate.

### **INTRODUCTION**

Valproic acid (VPA; 2-*n*-propylpentanoic acid) is a simple branched-chain fatty acid that is known worldwide for its anticonvulsant properties. It is mostly used for the treatment of several ty pes of se izures, b ipolar d isorders, v arious p sychiatric syndromes and m igraine (Peterson and Naunton, 2005; Perucca, 2002; Bialer and Yagen, 2007). In addition, VPA has recently emerged as a d rug t hat shows p otential in c ancert reatment (D uenas-Gonzalez, 2008). Although VPA has a broad range of clinical use, it is associated with several adverse effects. Hepatotoxicity is a well recognized complication of VPA therapy (G erber et al., 1979; B issellet al., 2001; Sztajnkrycer, 2 002). K nowledge of the pathophysiological mechanisms underlying this he patotoxicity is incomplete. Inhibition of mitochondrial catabolic pathways, for instance fatty acid β-oxidation, has strongly been implicated in the hepatotoxicity of VPA (Silva et al., 2008; Bjornsson, 2008; Silva et al., 2001a). In patients receiving valproate, an increase in serum levels of the branched-chain amino acids (BCAAs; leucine, i soleucine, and valine) and their intermediates as well as an increase in urinary excretion of these amino acids have been reported (Silva et al., 2001a; Anderson et al., 1994). These findings suggest that valproate adversely affects BCAAs metabolism.

VPA and the in termediates in the de gradation of the two B CAAs, isoleucine and valine undergo  $\beta$ -oxidation (Silva et al., 2002) generating structurally similar metabolites which suggest that VPA might use key enzymes of the B CAAs catabolic pathways for its own oxidation (figure 1). I sovaleryl-CoA, 2- methylbutyryl-CoA, and is obutyryl-CoA, the intermediates in BCAAs catabolism, are converted to 3-methylcrotonyl-CoA, tiglyl-CoA, and methacrylyl-CoA by i sovaleryl-CoA de hydrogenase (IVD, EC 1. 3.99.10), short branched-chain acyl-CoA dehydrogenase (SBCAD, EC 1. 3.99.12), and is obutyryl-CoA dehydrogenase (IBD, EC 1. 3.99.3), r espectively. These enzymes a rem embers of the acyl-CoA dehydrogenase f amily (ACD), which c onsists of ho mologous mitochondrial

flavoproteins that catalyze the  $\alpha$ , $\beta$ -dehydrogenation of a cyl-CoA thioester substrates to the corresponding *trans*-2-enoyl-CoAs (Battaile et al., 2004). These enzymes share many similar molecular and catalytic properties but differ with respect to their substrate specificities regarding length and configuration of the carbon backbone of their substrates.

Previous work from Li et al. (Li et al., 1991), Ito et al. (Ito et al., 1990) and our group (Silva et al., 2002; Silva et al., 2001b; Silva et al., 2004), has led to the partial resolution of the mitochondrial β-oxidation pathway of VPA. It was concluded that the enzymes involved in the oxidation of s traight-chain f atty acids i neluding ve ry l ong-, l ong-, m edium- a nd short-chain acyl-CoA dehydrogenases are not involved in the first dehydrogenation reaction of VPA. Ito et al. (Ito et al., 1990) originally showed that the rat 2-methyl-branched-chain acyl-CoA dehydrogenase was active with valproyl-CoA (VP-CoA) as substrate, but Willard et al (Willard et al., 1996) subsequently demonstrated that the human homologue was much less active with VP-CoA than its rat counterpart. In the current study we have investigated the involvement of the three BCAA acyl-CoA dehydrogenases with respect to the first step of the  $\beta$ -oxidation of VPA. We report the kinetic characterization of the SBCAD reaction with VP-CoA as substrate using the human purified enzyme. We also demonstrate that IVD is able to p roduce  $\Delta^{2(E)}$ -valproyl-CoA ( $\Delta^{2(E)}$ -VP-CoA), although with a much lower c atalytic efficiency. Furthermore, we have investigated the potential inhibitory effect of VPA and some of its mitochondrial metabolites especially in the form of the corresponding acyl-CoA intermediates. VP-CoA and valprovl-dephosphoCoA (VP-DephCoA) (Silva et al., 2001b: Silva et al., 2004), on the activity of IVD, IBD and SBCA D. The implications of these findings with respect to the treatment of patients by VPA are discussed.

MATERIAL AND METHODS

**Materials** 

Valproic a cid, hu man and bo vine se rum a lbumin, bicinchoninic a cid, ferrocenium

hexafluorophosphate, FAD, isovaleryl-CoA and isobutyryl-CoA were obtained from Sigma

Chemical Co. (St. Louis, MO). Tris was obtained from Merck (Darmstadt, Germany).

Heterologously expressed IVD, IBD and SBCAD were obtained as described before (Mohsen

and Vockley, 1995; Nguyen et al., 2002; Gibson et al., 2000). VP-CoA (Silva et al., 2001b),

VP-DephCoA (Silva et al., 2004), and 2-methylbutyryl-CoA (Rasmussen et al., 1990) were

synthesized as described previously.

Enzymatic activity measurement of the branched-chain acyl-CoA dehydrogenases

Experimental conditions were optimized for each enzyme in terms of protein, time and pH.

The reaction mixture contained 200 mM Tris·HCl pH 8.0, 50 µM FAD, 400 µM ferrocenium

hexafluorophosphate, 0.1 m g/mL bo vine serum a lbumin a nd su bstrate ( isovaleryl-CoA,

isobutyryl-CoA, 2-methylbutyryl-CoA) or VP-CoA. This mixture was added to an enzyme

solution containing 0.1 mg/mL BSA. Incubations were carried out at 37°C for variable time

periods depending on the enzyme and the substrate used. The reactions were terminated by

adding 10 µL 2 M HCl and afterwards the samples were placed on ice. After neutralization

with (2 M KOH)/(1 M MES pH 6.0), 10 μL 10 mM L-cysteine and 30 μL m ethanol were

added to the m ixture. L -cysteine i s a dded in o rder to r educe o xidized f errocinium

hexafluorophosphate which may interfere with the chromatographic separation. The samples

were c entrifuged at 2 0,000 x g f or 5 min and t he m etabolites in the supernatants we re

analysed by HPLC.

6

### Sample analysis by HPLC

Acyl-CoA esters were quantified by HPLC. Separation was performed at room temperature with a Perkin Elmer pump (PE series 200) and a Gilson 234 auto-sampling injector. A frit C-402X (Upchurch scientific), a 4.6 mm x 250 mm Supelcosil LC-18-DB (5  $\mu$ M) column (Supelco) and a guard column (4.6 mm x 20 mm) filled with the same packing material, were used. For gradient elution of b ranched-chain acyl-CoAs a b inary system of met hanol and 50 mM potassium phosphate pH 5.3 was used, whereas for the analysis of VPA intermediary metabolites a system of acet onitrile and 17 m M sodium p hosphate pH 6.9 was used. Acyl-CoA esters were detected with a UV detector (SPD-10A VP UV-VIS, Shimadzu) at 260 nm.

## Identification of $\Delta^2$ -VP-CoA by HPLC-ESI-MS/MS

HPLC-ESI-MS/MS analysis was performed using a triple-quadrupole TSQ Quantum HPLC tandem m ass spectrometer (MS/MS) from Thermo F innigan in the negative e lectrospray ionization (ESI) mode. The samples were injected onto an YMC-Pack Pro C<sub>4</sub> column (2.1 mm x 10 0 m m, YMC E urope GMBH) u sing a HPLC system c onsisting of a S urveyor MS-pump with degasser, a Surveyor autosampler and a column oven. The flow rate was set at 250  $\mu$ l/min. Elution of acyl-CoAs was achieved with tertiary system using solvent A (50 mM ammoniumacetate, pH 7.0), solvent B (1 00% a cetonitrile) and solvent C (20 m M ammoniumbicarbonate).

Separation was performed at 40°C and nitrogen was used as nebulizing gas while argon was used as collision gas at a pressure of 1.5 mTorr. The capillary voltage was 2.5 kV and the respective temperature was 350°C, with optimal collision energy of 30 eV. Acyl-CoA's were measured using multiple reaction monitoring (MRM) in the negative ionization mode, using

the transitions: m/z 445.5  $\rightarrow$  79.0 for C<sub>8</sub>-CoA (VP-CoA) and m/z 444.5  $\rightarrow$  79.0 for C<sub>8:1</sub>-CoA ( $\Delta^2$ -VP-CoA). The system was controlled by Xcalibur Software 2.0.

### Data analysis

The characterization of IVD, SBCAD a ctivities in the absence and presence of different inhibitors was performed by plotting the measured reaction rates as function of the substrate concentration (at a fixed concentration of the remaining components of the reaction). The steady state kinetic data of IVD, IBD and SBCAD activities were determined by nonlinear regression analysis using the SigmaPlot® 10.0 Technical Graphing Software (Systat, Inc.) and the Enzyme K inetics Module (v1.3). The Michaelis-Menten equation was used to calculate k inetic parameters ( $K_m$  and  $V_{max}$ ) of the enzyme reaction using the respective substrates. The inhibition constant  $K_i$  was calculated by nonlinear regression of the respective inhibition curves, using the mentioned software.

### **RESULTS**

#### Effects of VP-CoA and VP-DephCoA on the activity of IVD, IBD and SBCAD

Kinetic studies were performed with the three heterologously expressed and purified acyl-CoA dehydrogenases IVD, IBD and SBCAD. The corresponding activity was determined with their natural substrates, is ovaleryl-CoA, isobutyryl-CoA and 2-methylbutyryl-CoA, respectively. The kinetic constants  $K_m$  and  $V_{max}$  were calculated and are summarized in table 1.

The effect of VP-CoA and VP-DephCoA was tested on the three enzymes. The activities of IVD and SBCAD as measured at 0.2 mM isovaleryl-CoA and 0.2 mM 2-methylbutyryl-CoA, were decreased by about 75% and 45% respectively, at 1 mM VP-CoA. After incubation with 1 mM VP-DephCoA, both enzyme activities were decreased about 45%. The activity of IBD was not affected in the presence of VP-CoA and VP-DephCoA. I n order to further characterize the inhibition mechanism, the activity of IVD and SBCAD was determined as a function of the inhibitor concentration.

The ob tained r esults we re a nalysed by three lin earization methods of the Michaelis-Menten equation namely, Lineweaver-Burk, Eadie Hofstee and Hanes-Woolf plots. Analysis of the data pointed to a purely competitive mechanism of inhibition of both VP-CoA and VP-DephCoA with respect to IVD, as depicted in figure 2. However, with respect to SBCAD, these CoA esters appeared to have different inhibitory mechanisms. Indeed, whereas VP-CoA was found to inhibit SBCAD by a purely competitive mechanism, V P-DephCoA appeared to be a mixed-type inhibitor, as shown in figure 3. The Lineweaver-Burk Linearization plots and corresponding Dixon plots (Dixon, 1953) are exhibited in parallel, either in figure 2 or figure 3.

Assuming a purely competitive or mixed mechanism of inhibition, the inhibition constant ( $K_i$ ) was calculated. With respect to IVD activity, a  $K_i$  value of 74 ± 4  $\mu$ M was estimated for VP-CoA and of 170 ± 12  $\mu$ M f or VP -DephCoA. The inhibition constant ( $K_i$ ) of VP-CoA and V P-DephCoA for the activity of SBCAD was of 249 ± 29  $\mu$ M and 511 ± 96  $\mu$ M, respectively.

Involvement of branched-chain acyl-CoA dehydrogenases in the dehydrogenation of

VP-CoA

In order to identify the metabolizing enzyme of the first dehydrogenation step of the oxidative metabolism of VPA, the purified enzymes (IVD, IBD and SBCAD) were incubated with 1 mM

of VP -CoA. As s hown in figure 4,  $\Delta^{2(E)}$ -VP-CoA was n ot o bserved using I BD as enzyme

(Fig. 4-B), suggesting that IBD is not involved in the metabolism of VPA. However, both IVD

(Fig. 4-A) and SBCAD (Fig. 4-C) showed activity with VP-CoA as a substrate, amounting to

0.13 nmol/mg protein·min and 1.20 nmol/mg protein·min, respectively. Even though S BCAD

dehydrogenates VP-CoA a ta m uch lower rate than its natural s ubstrate, i.e.

2-methylbutyryl-CoA, the kinetic parameters were determined ( $K_{\rm m} = 304 \pm 51~\mu{\rm M}$  and  $V_{\rm max} =$ 

 $0.27 \pm 0.02$  nmol/mg protein·min).

An extra incubation was performed with SBCAD to identify the product of VP-CoA. Figure 5-B

shows the HPLC chromatograms related with the synthesis of  $\Delta^{2(E)}$ -VP-CoA, where SBCAD was

incubated for 1 hour with VP-CoA. In figure 5-A the same incubation was performed for 0 hour

and c onsequently there is n o p roduct f ormation. The same samples were analysed by

HPLC-ESI-MS/MS using multiple reaction monitoring in the negative ionization mode. Figure

5-C and 5-D shows the mass spectra of singly charged ions which are formed more abundantly.

The mass of a C<sub>8.1</sub>-CoA (889.8) was detected in samples where SBCAD was incubated for 1 hour

with VP-CoA (Fig. 5-D). This mass corresponds to  $\Delta^{2(E)}$ -VP-CoA, the dehydrogenation product

of VP-CoA.

**DISCUSSION** 

This study shows that valproate interferes with enzymes involved in the oxidative metabolism of

leucine and isoleucine. Specifically, two branched-chain acyl-CoA de hydrogenases (BCADs),

IVD and SBCAD, were found to participate in the oxidation of VPA. We have shown that human IVD is able to convert VP-CoA into  $\Delta^{2(E)}$ -VP-CoA, although at a lower rate than with its natural substrate, isovaleryl-CoA. VP-CoA has previously been shown to be oxidized at a significant rate by SBCAD from rat liver mitochondria (Ito et al., 1990) and by b acterially expressed rat and human SBCAD (Willard et al., 1996). We have used human SBCAD expressed in *E. coli* (Willard et al., 1996) to confirm and characterize the active role of SBCAD in the  $\beta$ -oxidation of VPA. These results are in agreement with the inhibition studies performed with VP-CoA and VP-DephCoA and the BCADs. Both VP A metabolites inhibit IVD by a purely competitive mechanism. IBD activity was not a ffected by these VP A metabolites. However, VP-CoA did inhibit SBCAD activity by a purely competitive mechanism whereas VP-DephCoA inhibited SBCAD activity through a distinct mechanism.

The observed inhibitory effects of the VPA metabolites as tested in this study, on the activity of the BCA Ds are most p robably due to structural s imilarities b etween the substrates of these enzymes and the valproate metabolites VP-CoA and VP-DephCoA. IVD oxidizes  $\beta$ -branched acyl-CoAs such as isovaleryl-CoA, but both SBCAD and IBD oxidize short-chain acyl-CoAs with a branched  $\alpha$ -position (figure 6). Due to its  $\alpha$ -branched configuration, VP-CoA would have appeared to be a good substrate analogue for both SBCAD and IBD. However, VP-CoA is a substrate analogue for SBCAD but not for IBD. The natural substrate of IBD is isobutyryl-CoA, which has a smaller acyl moiety than the VP-CoA acyl moiety, hence the apparent lack of interaction between VP-CoA and IBD is probably because of the limited size of the isobutyryl moiety binding pocket (Battaile et al., 2004). SBCAD is able to oxidize  $\alpha$ -branched acyl-CoAs such as 2-methyl-butyryl-CoA and straight chain substrates as butyryl-CoA and hexanoyl-CoA (He et a l., 2003), and therefore its active site seems to be the best ac commodating for the dehydrogenation of VP-CoA.

Unexpectedly, VP-CoA was found to be oxidized by IVD. Although the reaction occurs at a very

low rate, it is still rather surprising since this enzyme handles  $\beta$ - and not  $\alpha$ -branched acyl-CoAs.

IVD has a larger binding pocket than IBD (Battaile et al., 2004), which apparently allows the

enzyme to act upon VP-CoA with its five carbon backbone. This result is in agreement with the

competitive mechanism of inhibition seen for IVD when VP-CoA was used as an inhibitor.

Since SBCAD dehydrogenates VP-CoA to  $\Delta^{2(E)}$ -VP-CoA and is significantly inhibited by both

VP-CoA and V P-DephCoA, it would be expected that patients under VPA therapy would

accumulate the en dogenous substrate of SBCAD, 2-methylbutyryl-CoA. However, no studies

have reported the increase of metabolites derived from 2 -methylbutyryl-CoA perhaps because

IBD also has activity towards this substrate (Nguyen et al., 2002) substituting, at least in part,

SBCAD activity in case of its malfunction or inhibition.

It has been shown that a dysregulated BCAA metabolism makes an independent contribution to

development of insulin resistance and glucose intolerance in obe se humans (Newgard et al.,

2009). Insulin resistance associated to weight gain has also been reported during VPA treatment

(Masuccio e t a l., 2010; Ve rrotti e t a l., 2010). However, i t is un known t o what e xtent the

interference of v alproate on the BCAA o xidation shown in this paper is r elated with the

significant we ight ga in p otentially a ssociated with VP A. It r emains to be e stablished if the

inhibitory e ffects, as o bserved in this study, are en ough to a count for the well-recognized

potential liver toxicity of VPA. In this respect, it is important to mention that although there are a

significant number of patients suffering from side effects, VPA-induced hepatotoxicity seems to

be idiosyncratic and only affects a small group of patients.

In summary, we have shown that VPA can interfere with the activity of some enzymes from the

ACD family, n amely, IVD and S BCAD. Ta king in to a count that AC Ds have overlapping

activity with different substrates (B attaile et al., 2004; T iffany et al., 1997), the drug might

exacerbate the impairment of genetically affected routes and elicit a toxic condition (Silva et al.,

DMD Fast Forward. Published on March 23, 2011 as DOI: 10.1124/dmd.110.037606 This article has not been copyedited and formatted. The final version may differ from this version.

DMD # 37606

2008). In fact, it has been suggested already that the intake of VPA should be avoided in patients

with inborn errors affecting mitochondrial metabolism (Silva et al., 2008). For that reason, we

conclude that VPA administration should be a voided in cases of inborn deficiencies affecting

certain A CDs or a ffecting the leucine and isoleucine oxi dative p athways (Vockley a nd

Ensenauer, 2006; Korman, 2006).

**ACKNOWLEDGEMENTS** 

We thank Arno van Cruchten for his expert technical assistance.

**AUTHORSHIP CONTRIBUTIONS** 

Participated in research design: Luís PB, Ruiter J, IJlst L, Duran M, Wanders RJ and Silva MF

Conducted experiments: Luís PB

Contributed new reagents or analytical tools: Mohsen A-W and Vockley J

Performed data analysis: Luís PB, Ruiter J, IJlst L and Silva MF

Wrote or contributed to the writing of the manuscript: Luís PB, IJlst L, Almeida IT, Duran M,

Vockley J, Wanders RJ and Silva MF

Acquired funding for the research: Wanders RJ and Silva MF

13

### **REFERENCES**

- Anderson GD, A cheampong AA and Levy RH. (1994) Interaction between valproate and branched-chain amino acid metabolism. *Neurology* **44**:742-744.
- Battaile KP, N guyen TV, Vo ckley J and K im JJ. (2004) Structures of iso butyryl-CoA dehydrogenase and enzyme-product complex: comparison with isovaleryl- and short-chain acyl-CoA dehydrogenases. *J Biol Chem* **279**:16526-16534.
- Bialer M and Yagen B. (2007) Valproic Acid: second generation. *Neurotherapeutics* **4**:130-137.
- Bissell DM, Go res GJ, Laskin DL and Hoofnagle JH. (2001) Dr ug-induced liver in jury: mechanisms and test systems. *Hepatology* **33**:1009-1013.
- Bjornsson E. (2008) He patotoxicity associated with antiepileptic drugs. *Acta Neurol Scand* **118(5)**:281-290.
- Dixon M. (1953) The determination of enzyme inhibitor constants. *Biochem J* 55:170-171.
- Duenas-Gonzalez A, Candelaria M, Perez-Plascencia C, Perez-Cardenas E, de la Cruz-Hernandez E and Herrera LA. (2008) Valproic acid as epigenetic cancer drug: preclinical, clinical and transcriptional effects on solid tumors. *Cancer Treat Rev* **34**:206-222.
- Gerber N, Dickinson RG, Harland RC, Lynn R K, Houghton D, Antonias JI, Schimschock JC. (1979) Reye-like syndrome associated with valproic acid therapy. *J Pediatr* **95**:142-144.
- Gibson KM, Burlingame TG, Hogema B, Jakobs C, Schutgens RB, Millington D, Roe CR, Roe DS, Sweetman L, Steiner RD, Linck L, Pohowalla P, Sacks M, Kiss D, Rinaldo P, Vockley J. (2000) 2-Methylbutyryl-coenzyme A dehydrogenase deficiency: a new inborn error of L-isoleucine metabolism. *Pediatr Res* **47**:830-833.

- He M., Burghardt TP and Vockley J. (2003) A novel approach to the characterization of substrate s pecificity in s hort/branched ch ain Acyl-CoA dehydrogenase. *J Biol Chem* **278**:37974-37986.
- Ito M., Ikeda Y, Arnez JG, Finocchiaro G and Tanaka K. (1990) The enzymatic basis for the metabolism and inhibitory effects of valproic acid: dehydrogenation of valproyl-CoA by 2-methyl-branched-chain acyl-CoA dehydrogenase. *Biochim Biophys Acta* **1034**:213-218.
- Korman S H . (2006) Inborn errors of isoleucine degradation: a rev iew. *Mol Genet Metab* **89**:289-299.
- Li J, Norwood DL, Mao LF and Schulz H. (1991) Mitochondrial metabolism of valproic acid. *Biochemistry* **30**:388-394.
- Masuccio F, Verroti A, Chiavaroli V, Giorgis T, Giannini C, Chiarelli F, and Mohn A (2010)

  Weight gain and insulin resistance in children treated with valproate: the influence of time. *J Child Neurol* 25(8) 941-947
- Mohsen AW and Vockley J. (1995) High-level expression of an altered c DNA encoding human isovaleryl-CoA dehydrogenase in Escherichia coli. *Gene* **160**:263-267.
- Newgard CB, An J, Bain JR, Muehlbauer MJ, Stevens RD, Lien LF, Haqq AM, Shah SH, Arlotto M, Slentz CA, Rochon J, Gallup D, Ilkayeva O, Wenner BR, Yancy WS, Eisenson H, Musante G, Surwit RS, Millington DS, Butler MD, and Svetkey LP (2009) A branched-chain amino a cid-related metabolic signature that differentiates obese and lean humans and contributes to insulin resistance. *Cell Metab* **9**, 311-326.
- Nguyen TV, Andresen BS, Corydon TJ, Ghisla S, Abd-El R azik N, Mohsen AW, et al. (2002) Identification of isobutyryl-CoA dehydrogenase and its deficiency in humans. *Mol Genet Metab* 77:68-79.
- Perucca E. (2002) Pharmacological and therapeutic properties of valproate: a summary after 35 years of clinical experience. *CNS Drugs* **16**: 695-714.

- Peterson G M and Naunton M. (2005) Valproate: a simple chemical with so much to offer. *J Clin Pharm Ther* **30**: 417-421.
- Rasmussen JT, Borchers T and Knudsen J. (1990) Comparison of the binding affinities of acyl-CoA-binding protein and fatty-acid-binding protein for long-chain acyl-CoA esters. Biochem J 265:849-855.
- Silva M F, R uiter JP, Ijlst L, J akobs C, D uran M, de Alm eida I T, W anders R J (2001a) Differential effect of valproate and its Delta2- and Delta4-unsaturated metabolites, on the beta-oxidation r ate o f long-chain a nd medium-chain f atty a cids. *Chem Biol Interact* 137:203-212.
- Silva MF, Ruiter JP, Ijlst L, Allers P, ten Brink HJ, Jakobs C, Duran M, Tavares de Almeida I, Wanders RJ. (2001b) Synthesis and intramitochondrial levels of valproyl-coenzyme A metabolites. *Anal Biochem* **290**:60-67.
- Silva M F, R uiter JP, Ove rmars H, Bootsma AH, va n Gennip AH, Jakobs C, D uran M, Tavares de Almeida I, Wanders RJ. (2002) Complete beta-oxidation of valproate: cleavage of 3-oxovalproyl-CoA by a mitochondrial 3-oxoacyl-CoA thiolase. *Biochem J* **362**:755-760.
- Silva M F, I jlst L, Al lers P, Jakobs C, Du ran M, de Alm eida I T, W anders RJ (2004) Valproyl-dephosphoCoA: a n ovel m etabolite of valproate f ormed in vit ro i n r at li ver mitochondria. *Drug Metab Dispos* **32**:1304-1310.
- Silva MF, Aires CC, Luis PB, Ruiter JP, Ijlst L, Duran M, Wanders RJ, Tavares de Almeida
  I. (2008) Valproic acid metabolism and its effects on mitochondrial fatty acid oxidation:
  A review. *J Inherit Metab Dis* 31:205-216.
- Sztajnkrycer MD. (2002) Valproic acid toxicity: overview and management. *J Toxicol Clin Toxicol* **40**:789-801.

- Tiffany KA, Roberts DL, Wang M, Paschke R, Mohsen AW, Vockley J and Kim JJ. (1997)

  Structure of human isovaleryl-CoA dehydrogenase at 2.6 A resolution: structural basis for substrate specificity. *Biochemistry* **36**:8455-8464.
- Verrotti A, D'Egidio C, Mohn A, Coppola G, and Chiarelli (2010) Weight gain following treatment with valproic acid: pathogenic mechanisms and clinical implications. *Obes Rev* 1467-789.
- Vockley J and E nsenauer R. (2006) I sovaleric a cidemia: ne w a spects of g enetic and phenotypic heterogeneity. *Am J Med Genet C Semin Med Genet* **142**:95-103.
- Willard J, Vicanek C, Battaile KP, Van Veldhoven PP, Fauq AH, Rozen R and Vockley J. (1996) C loning of a cDNA for s hort/branched c hain a cyl-Coenzyme A dehydrogenase from rat and characterization of its tissue ex pression and s ubstrate s pecificity. *Arch Biochem Biophys* **331**:127-133.

Footnotes:

This work was supported by Fundação para a Ciência e a Tecnologia, Lisboa, Portugal [Grant

SFRH/BD/25913/2005] and Jerry Vockley was supported in part by NIH [Grant R01-DK54936].

-----

Part of this work was presented as an oral communication at SSIEM (Society for the Study of

Inborn Errors of Metabolism) – Annual Symposium 2008, Lisbon, Portugal. Abstract published

as: Luís PBM, Ruiter JP, Ijlst L, Ofman R, Diogo L, Garcia P, Duran M, Vockley J, Tavares de

Almeida I, Wanders RJA and Silva MFB, Interference of Valproic Acid on the Branched Chain

Amino Acid Oxidative Metabolism, J Inherit Metab Dis 31 (Suppl 1):7, 2008.

-----

R.J.A.W. and M.F.B.S. are equal last authors.

# **LEGENDS FOR FIGURES:**

**Figure 1** – Schematic representation of the catabolic pathways of the branched-chain amino acids (leucine, isoleucine and valine) and valproic acid, highlighting the reactions of the first cycle of β-oxidation and corresponding enzymes of the initial dehydrogenation (1- I VD: Isovaleryl-CoA dehydrogenase; 2- SBCAD: Short branched chain acyl-CoA dehydrogenase; 3- IBD: Isobutyryl-CoA dehydrogenase)

**Figure 2** – I nhibitory e ffect of V P-CoA and V P-DephCoA on the activity of heterologously expressed human isovaleryl-CoA dehydrogenase (IVD). Lineweaver Burk linearization plots of IVD activity with isovaleryl-CoA as a substrate in the presence of VP-CoA (**A**) and VP-DephCoA (**B**). Dixon plots of IVD activity in the presence of VP-CoA (**C**) and VP-DephCoA (**D**). Incubations were carried at 37°C, pH 8, for 10 min.

**Figure 3** – I nhibitory e ffect of V P-CoA and V P-DephCoA on the activity of heterologously expressed human short branched-chain acyl-CoA dehydrogenase (SBCAD). Lineweaver Burk linearization plots of SBCAD activity with 2-methylbutyryl-CoA as a substrate in the presence of VP-CoA (**A**) and VP-DephCoA (**B**). Dixon plots of SBCAD activity in the presence of VP-CoA (**C**) and VP-DephCoA (**D**). Incubations were carried at 37°C, pH 8, for 30 min.

**Figure 4** – Activity of the br anched-chain a cyl-CoA de hydrogenases ( IVD, IBD a nd SBCAD) using VP-CoA as a substrate. HPLC analysis of the incubation mixtures with IVD (**A**), IBD (**B**) and SBCAD (**C**) using 1 mM VP-CoA as substrate. Incubations were carried out at 37°C for 30 min (——) and 0 min (---). Peaks: 1: VP-CoA and 2:  $\Delta^{2(E)}$ -VP-CoA.

**Figure 5** – Identification of the oxidation product of VP-CoA using SBCAD. (A) and (B): HPLC chromatograms of incubations of SBCAD with VP-CoA, carried out at 37° for 0h or 1h, respectively. (C) and (D): MS spectra (singly charged ions) of incubations of SBCAD with VP-CoA, carried out at 37° for 0h or 1h, respectively. Metabolites: 1: VP-CoA, 2:  $\Delta^{2(E)}$ -VP-CoA.

**Figure 6** – Chemical structures of i sovaleryl-CoA, 2- methylbutyryl-CoA and isobutyryl-CoA, intermediates from the leucine, isoleucine and valine pathways, respectively, and VP-CoA, the CoA thioester of valproic acid.

**Table 1** – Kinetic parameters ( $K_{\rm m}$  and  $V_{\rm max}$ ) of IVD, IBD and SBCAD activities using the substrates, isovaleryl-CoA, iso butyryl-CoA and 2-methylbutyryl-CoA, r espectively. Incubations were carried at 37°C, pH 8 for 10 min for IVD assay and for 30 min for IBD and SBCAD assays. (The results are the mean +/- SD from two to three independent experiments).

Enzyme	Substrate	$K_{\rm m} (\mu { m M})$	$V_{\rm max}$ (µmol/(mg protein·min))
IVD	Isovaleryl-CoA	$125 \pm 2.3$	$77 \pm 6.7$
IBD	Isobutyryl-CoA 24	$\pm 0.6$	$32 \pm 1.5$
SBCAD	2-Methylbutyryl-CoA	$12 \pm 2.0$	$12 \pm 0.2$

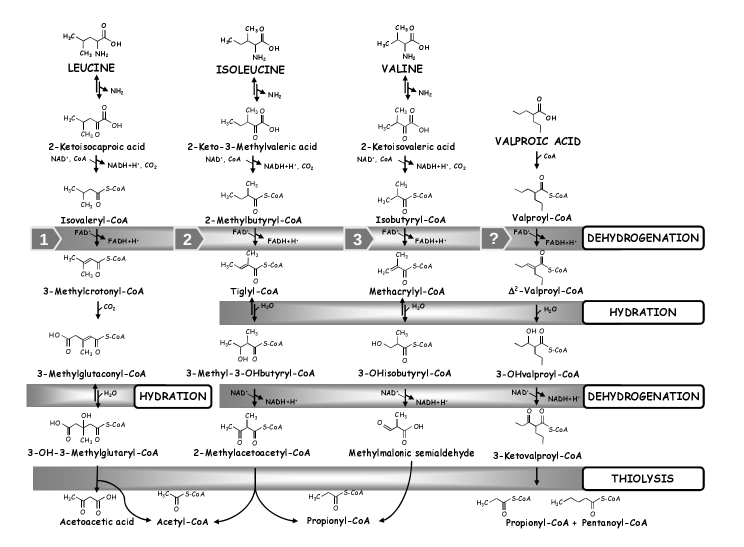
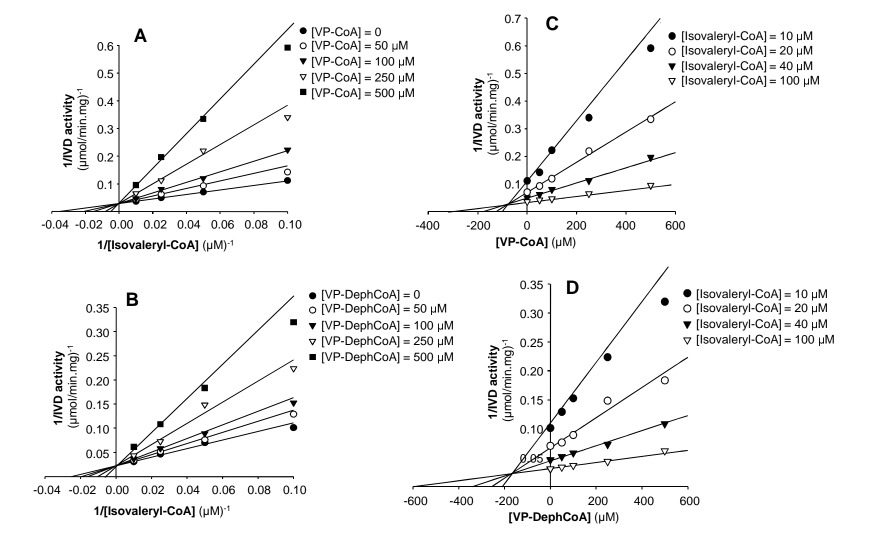


Figure 1

Figure 2



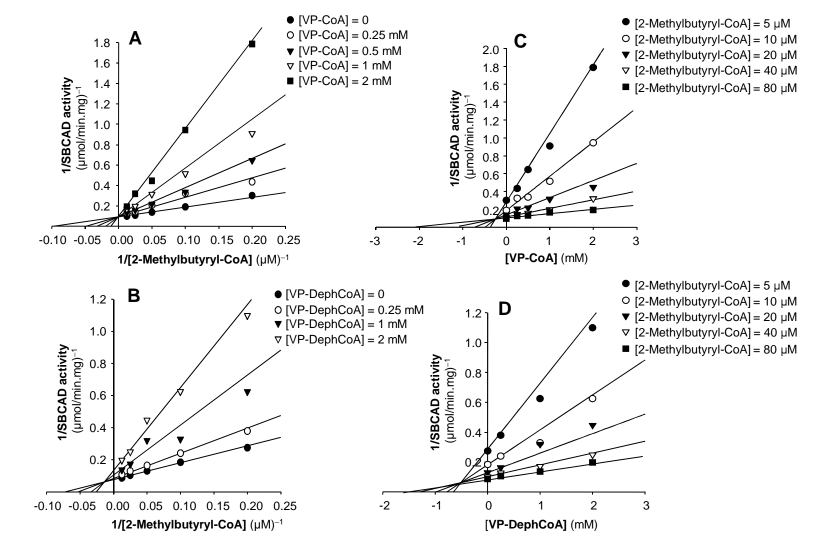


Figure 3

Figure 4

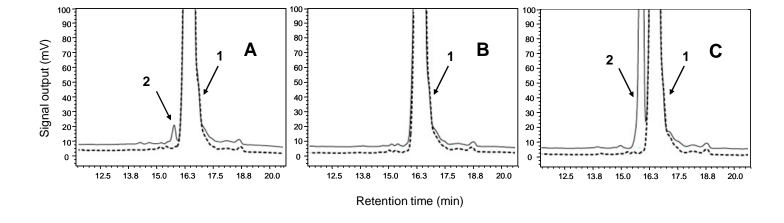
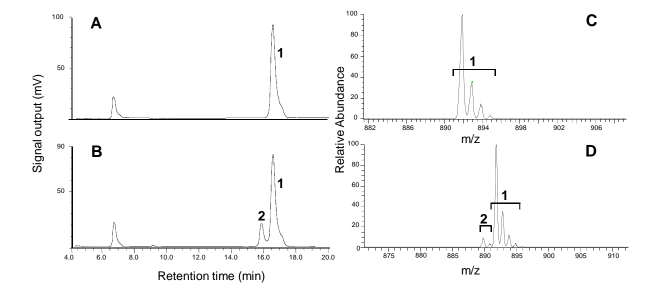


Figure 5



Isovaleryl-CoA

2-Methylbutyryl-CoA

H<sub>3</sub>C 
$$\alpha$$
 SCoA

Isobutyryl-CoA

Valproyl-CoA