

## <sup>1</sup> Objectives:

# 1 Neuroprotective effects of tibolone 2 during astrocytic metabolic 3 inflammation: a network based 4 approach 5

## 6 Abstract:

### 7 1.1. Introduction

#### 8 Astrocyte-Neuron Metabolic Relationships

9 Astrocytes are the most abundant cells in the human brain and play important roles in  
10 the central nervous system (CNS) [1]. They are highly associated to several homeostatic fun-  
11 ctions such as glutamate, ion, and water homeostasis, energy storage in the form of glycogen,  
12 synapse formation and remodeling, defense against oxidative stress, scar formation, tissue  
13 repair and modulation of synaptic activity via the release of gliotransmitters [2]. Astrocytes  
14 metabolize glucose in anaerobic way to produce lactate, which is released to neurons through  
15 monocarboxylate transporters [3]. Lactate is used in neurons as an energy substrate after  
16 its conversion to pyruvate and subsequently to ATP via oxidative phosphorylation [4]. As-  
17 trocytes play an important role in glutamate mediated synaptic activity [5]; according to the  
18 astrocyte–neuron lactate shuttle model, astrocytes respond to glutamate induced activation  
19 by increasing their rate of glucose uptake and the release of lactate into the extracellular spa-  
20 ce, increasing the lactate available to be used by neurons to supply their energetic needs [6].  
21 Glutamate is uptaken by astrocytes through the glutamate aspartate transporter and glial  
22 glutamate transporter-1, inducing events that involves the activation of  $\text{Na}^+/\text{K}^+$ -ATPase  
23 and maintaining extracellular glutamate at homeostatic levels [7]. Part of incorporated glu-  
24 tamate is converted to glutamine through glutamine synthetase, which is only associated to  
25 glial cells and released to neurons using electroneutral systems-N transporters coupled to  
26  $\text{Na}^+$  and  $\text{H}^+$  [8]. In neurons glutaminase enzyme converts glutamine back into glutamate  
27 which can be used again for neurotransmission or metabolized into the neuronal Krebs cycle  
28 [9]. Astrocytes release many other substances related to synaptic transmission [10]. However  
29 D-serine, a neurotransmitter that act as a coagonist with glutamate at NMDA receptors is

one of the most important [5]. Due only glial cells can synthesize serine, all available D-serine at synapsis is associated to be primarily produced and secreted by astrocytes [8]. D-serine is synthesized in astrocytes by serine racemase from L-serine [11]. Additionally to these energy and synaptic support associated functions, astrocytes also play an important role in the reduced glutathione (GSH) metabolism of the brain [12]. GSH is the major cellular antioxidant and plays an important neuroprotective role [13]. Cellular GSH levels are closely correlated with cell survival under adverse conditions [14]. GSH is synthesized from glutamate, cysteine, and glycine and release directly from Astrocytes through GSH transporters ion-independent, net transport is concentration-gradient dependent [15].

This strong metabolic cooperation between astrocytes and neurons allows to predict that even an small astrocytic dysfunction might cause and/or contribute neurodegenerative processes.

## Inflammation

Inflammation is a biological response to injury, metabolic disorders or infection and its dysregulation underlies many complex diseases. The inflammatory response plays an important role in the defense mechanism against any threat to normal integrity that is required for the maintenance of the healthy state. Neuroinflammation in CNS is mediated by glial cells that acquire reactive phenotypes to participate in neuronal repair mechanisms. Inflammatory response induce changes in glucose metabolism and release of proinflammatory factors. As glial cells, astrocytes are highly sensitive cells to inflammatory mediators, they respond to inflammation through a complex reaction named astrogliosis. Although glial cells execute several beneficial activities in the CNS, these same cell types also act as a source of inflammatory mediators and as generators of reactive oxidant species (ROS) that have the potential to damage neurons. Astrogliosis is also characterized by a faulty regulation of mitochondrial dynamics that result in defective mitochondria. Mitochondrial failure induces the deregulation of  $\text{Ca}^{2+}$  homeostasis and increased ROS generation, both of which are linked to neurotoxicity. At metabolic level, inflammatory process has been associated to an increase of free saturated fatty acid in comparison with healthy conditions in some brain tissues. The increase of free saturated fatty acid induce metabolic inflammation, a response associated with the induction of diverse intracellular stresses, such as mitochondrial oxidative stress, ER stress, and autophagy defects. Lipid excess in metabolic inflammation activates hypothalamic  $\text{IKK}\beta/\text{NF-}\kappa\beta$ , which ultimately impairs hypothalamic leptin and insulin signaling and further triggers the synthesis and release of increased amounts of ROS and proinflammatory cytokines, such as  $\text{TNF-}\alpha$  and IL-6, to contribute to central metabolic inflammation and to sustain the neuroinflammatory state. Enhanced ROS generation by reactive glial cells trigger mitochondria dysfunction, which causes the onset of neuronal death (apoptotic or necrotic), which is the prerequisite for a diverse number of neurodegenerative conditions.

## 67 SysBio Inflammation

## 68 Tibolone

## 69 1.2. Material and Methods

### 70 Tissue Specific Model Construction

71 The tissue specific model construction process started with the identification of all enzyme-  
72 coding genes expressed over the mean in at least 50 % of samples for healthy human astrocy-  
73 tes indexed in the GEO database [16] as GSE73721 [17]. Gene identifiers conversion from  
74 GeneCards[18] to ENTREZ [19] was performed through ‘UniProt.ws’ R Package [20]. Reac-  
75 tions associated with the identified genes were mapped from the Human Genome Scale Me-  
76 tabolic Reconstruction RECON 2.04 downloaded from the VMH Lab (<https://vmh.uni.lu>)  
77 [21]. The R package ‘g2f’ [22] was used to identify and fill the gaps using all no gene as-  
78 sociated reactions included in RECON 2.04, as well as to identify and remove all blocked  
79 reactions from the reconstruction. All reactions involved in the conversion of extracellular  
80 glutamate, glycine, cysteine and glucose to extracellular glutamine, glycine, serine-D, redu-  
81 ced glutathione, lactate and ATP respectively were added. Exchange reactions were limited  
82 to components of the Dulbecco’s Modified Eagle Medium (DMEM) as input and gliotrans-  
83 mitters (glutamine, D-serine, ATP, glutamate), reduced glutathione, lactate, glucose, nitric  
84 oxide, prostaglandins and leukotrienes as output. Finally, syntax, mass-charge validation and  
85 creation of SBML files were carried out through the ‘minval’ R Package [23]. Reaction limits  
86 (upper and lower bounds) were constrained proportional to the mean gene expression repor-  
87 ted for genes included in Gene-Protein-Reaction (GPR) [24] associated to each reaction in  
88 samples of 47 to 63 years old using ‘exp2flux’ R package [25]. All analysis were done by the  
89 ‘sybil’ [26] R Package running under R 3.3.1 [27].

### 90 Flux Balance Analysis

91 Flux Balance Analysis (FBA) is a linear optimization method for simulating metabolism that  
92 allows to identify the set of reactions involved in the production of a biological response within  
93 a metabolic model [28]. The metabolic reactions are represented internally as a stoichiometric  
94 matrix ( $S$ ), of size  $m \times n$ , where  $m$  represents the compounds and  $n$  the reactions; the entries  
95 in the matrix are the stoichiometric coefficients of the metabolites participating in a reaction  
96 [29]. The flux through all of the reactions in a network is represented by the vector  $v$ , which  
97 has a length of  $n$ . The concentrations of all metabolites are represented by the vector  $x$ , with  
98 length  $m$ . The systems of mass balance equations at steady state,  $\frac{d_x}{d_t} = 0$  or  $S * v = 0$ . FBA  
99 seeks to maximize or minimize an objective function which can be any linear combination  
100 fluxes, to obtain a flux for each reaction, indicating how much each reaction contributes  
101 to the objective function [28]. FBA for healthy, inflamed and medicated scenarios was

resolved using GLPK 4.60, setting the generic human biomass reaction included in RECON 2.04 and each one of reactions described in table **1-1** as objective functions. Models were analyzed by comparing fluxes between scenarios, metabolites production rate and sensitivity analysis.

**Table 1-1:** Main metabolic capabilities associated to astrocytes represented as the set of objective functions used to evaluate neuroprotective effects of Tibolone under inflamed scenarios

ID	FORMULA REACTION	DESCRIPTION
Glu2Gln	$1 \text{ glu\_L[e]} \Rightarrow 1 \text{ gln\_L[e]}$	Glutamate - Glutamine Cycle
Gly2SerD	$1 \text{ gly[e]} \Rightarrow 1 \text{ ser\_D[e]}$	Glycine to D-serine conversion
Glc2Lac	$1 \text{ glc\_D[e]} \Rightarrow 2 \text{ lac\_L[e]}$	Lactate production from Glucose
Glc2ATP	$1 \text{ glc\_D[e]} \Rightarrow 36 \text{ atp[e]}$	ATP production from Glucose
Cys2GTHRD	$1 \text{ cys\_L[e]} + 1 \text{ glu\_L[c]} + 1 \text{ gly[c]} \Rightarrow 1 \text{ gthrd[e]}$	Catch of Cysteine to produce reduced Glutathione

## Metabolic Scenarios

To test neuroprotective effects of tibolone during astrocytic metabolic inflammation we define three different metabolic scenarios. A ‘healthy’ scenario, where palmitate uptake rate was freely set by optimizer; an ‘inflamed’ scenario, where uptake rate of palmitate was forced to be stable in the mean of the half maximal inhibitory concentration (IC<sub>50</sub>) value for all objective functions included in table **1-1**. IC<sub>50</sub> values were calculated through a robustness analysis performed using uptake of palmitate (‘EX\_hdca(e)’ in RECON 2.04) as control reaction and a 1000 points in the range from 0 to 1 mMgDW<sup>-1</sup>h<sup>-1</sup> for each objective function. Uptake value where each objective function reached IC<sub>50</sub> was selected and subsequently averaged. Finally, a medicated scenario, defined as an inflamed scenario that include 279 reactions associated with tibolone and estradiol-derived compounds metabolism. Ten specific reactions described in table **1-2** associated to specific Tibolone action mechanism non included in RECON 2.04 were added to medicated scenario.

## Metabolic Changes

Metabolic changes across metabolic scenarios were measured through two different approximations. Flux differences for each reaction between optimized scenarios were measured using

**Table 1-2:** Set of reactions associated to tibolone specific action mechanism in brain reported by Kloosterboer, H. J. (2004) added to medicated scenario model.

ID	FORMULA REACTION	DESCRIPTION
T1	tibolone[e] $\Leftrightarrow$	Tibolone exchange reaction
T2	tibolone[e] $\Leftrightarrow$ a3OHTibolone[e]	3 $\alpha$ hydroxytibolone interconversion
T3	tibolone[e] $\Leftrightarrow$ b3OHTibolone[e]	3 $\beta$ hydroxytibolone interconversion
T4	tibolone[e] $\Rightarrow$ d4tibolone[e]	Tibolone $\Delta^4$ isomer formation
T5	b3OHTibolone[e] $\Rightarrow$ d4tibolone[e]	Tibolone $\Delta^4$ isomer formation from 3 $\beta$ -hydroxytibolone
T6	a3OHTibolone[e] $\Rightarrow$ estradiol[c]	Estradiol receptor agonist action mechanism of 3 $\alpha$ -hydroxytibolone
T7	b3OHTibolone[e] $\Rightarrow$ estradiol[c]	Estradiol receptor agonist action mechanism of 3 $\beta$ -hydroxytibolone
T8	d4tibolone[e] $\Rightarrow$ prgstrn[c] + tststerone[c]	Progesterone and androgen receptor activation by tibolone $\Delta^4$ isomer
T9	a3OHTibolone[e] $\Leftrightarrow$ a3SOTibolone[e]	3 $\alpha$ hydroxytibolone interconversion to sulfated inactive compounds
T10	a3SOTibolone[e] $\Rightarrow$	Tibolone inactive form in blood

the fold change calculated as described in equation 1-1.

$$foldChange = \frac{valueModel2 - valueModel1}{|valueModel1|} \tag{1-1}$$

120 Additionally, to obtain a full perspective about inflammation effects in metabolites pro-  
121 duction, the production of each metabolite was set as objective function in each metabolic  
122 scenario and differences were evaluated as well as flux differences.

123 **Proinflammatory, Antiinflammatory and Tibolone Action Mechanism**

124 **Associated Enzymes**

125 Identification of enzymes involved in proinflammatory and antiinflammatory responses as  
126 well as in the tibolone action mechanism were identified through several sensitivity analysis  
127 as follows: Proinflammatory enzymes, are those that catalyze reactions that being knocked  
128 out allows an increase of objective function value. Antiinflammatory enzymes, are those  
129 associated to reactions that being knocked out reduce even more the objective function  
130 value. Tibolone action mechanism associated enzymes are those that catalyze reactions that  
131 being knocked out inhibit entirely the metabolic effect of tibolone.

132 **1.3. Results**

133 **1.4. Conclusion**

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