

A mathematical model of the mevalonate cholesterol biosynthesis pathway

Article

Accepted Version

Creative Commons: Attribution-Noncommercial-No Derivative Works 4.0

Pool, F., Currie, R., Sweby, P. K., Salazar, J. D. and Tindall, M. J. (2018) A mathematical model of the mevalonate cholesterol biosynthesis pathway. Journal of Theoretical Biology, 443. pp. 157-176. ISSN 0022-5193 doi: https://doi.org/10.1016/j.jtbi.2017.12.023 Available at http://centaur.reading.ac.uk/74972/

It is advisable to refer to the publisher's version if you intend to cite from the work. See <u>Guidance on citing</u>.

To link to this article DOI: http://dx.doi.org/10.1016/j.jtbi.2017.12.023

Publisher: Elsevier

All outputs in CentAUR are protected by Intellectual Property Rights law, including copyright law. Copyright and IPR is retained by the creators or other copyright holders. Terms and conditions for use of this material are defined in the <u>End User Agreement</u>.

www.reading.ac.uk/centaur

CentAUR



Central Archive at the University of Reading

Reading's research outputs online

A mathematical model of the mevalonate cholesterol 1 biosynthesis pathway 2

Frances Pool^a, Richard Currie^b, Peter K. Sweby^c, José Domingo Salazar^d, Marcus J. Tindall^{c,e,*} 4

^aInstitute of Ophthalmology, University College London, Gower Street, London, United Kingdom, WC1E 6BT. ^bSyngenta, Jealott's Hill International Research Centre, Bracknell, Berkshire, United Kingdom, RG42 6EY.

^cDepartment of Mathematics and Statistics, University of Reading, Whiteknights, Reading, United Kingdom, RG6 6AX.

^dAstraZeneca, Unit 310, Cambridge Science Park, Milton Road, Cambridge, Cambridgeshire, United Kingdom, CB4 0FZ

^eInstitute of Cardiovascular and Metabolic Research, University of Reading, Whiteknights, Reading, United Kingdom, RG6 6AA.

Abstract 15

3

5

6

7

8

9

10

11

12

13

14

We formulate, parameterise and analyse a mathematical model of the mevalonate pathway, a key pathway in the synthesis of cholesterol. Of high clinical importance, the pathway incorporates rate limiting enzymatic reactions with multiple negative feedbacks. In this work we investigate the pathway dynamics and demonstrate that rate limiting steps and negative feedbacks within it act in concert to tightly regulate intracellular cholesterol levels. Formulated using the theory of nonlinear ordinary differential equations and parameterised in the context of a hepatocyte, the governing equations are analysed numerically and analytically. Sensitivity and mathematical analysis demonstrate the importance of the two rate limiting enzymes 3-hydroxy-3methylglutaryl-CoA reductase and squalene synthase in controlling the concentration of substrates within the pathway as well as that of cholesterol. The role of individual feedbacks, both global (between that of cholesterol and sterol regulatory element-binding protein 2; SREBP-2) and local internal (between substrates in the pathway) are investigated. We find that

Preprintesponding outharm Renn Austral due Bio Department of Mathematics and Statiss tics, University of Reading, Whiteknights, Reading, UK, RG6 6AX. Tel.: +44 118 378 8989.

Email address: m.tindall@reading.ac.uk (Marcus J. Tindall)

whilst the cholesterol SREBP-2 feedback regulates the overall system dynamics, local feedbacks activate within the pathway to tightly regulate the overall cellular cholesterol concentration. The network stability is analysed by constructing a reduced model of the fall pathway and is shown to exhibit one real, stable steady-state. We close by addressing the biological question as to how farnesyl-PP levels are affected by CYP51 inhibition, and demonstrate that the regulatory mechanisms within the network work in unison to ensure they remain bounded.

¹⁶ Keywords: nonlinear ordinary differential equation, feedback, HMGCR,

17 squalene synthase

18 **1. Introduction**

The mevalonate pathway is an important metabolic pathway present in all eukaryotes, fungi and some bacteria [6, 13]. It is responsible for many processes within the cell including biosynthesis of cholesterol, cell wall maintenance, hormone production, protein lipidation and anchoring and is part of steroid biosynthesis.

The body produces around 80% of cholesterol it needs [40]. A large percent-24 age of this is synthesised by the liver via a series of reactions. In mammalian 25 cells cholesterol is a substrate for a number of other reactions [6]. Over ac-26 cumulation of cholesterol can lead to cellular toxicity [18], whilst insufficient 27 cholesterol levels result in compromised cell structure and function. Thus 28 it is important that cholesterol levels are tightly regulated within the cell. 29 This is known as cellular cholesterol homeostasis and it works by balanc-30 ing the influx, utilisation and efflux of cholesterol to maintain intracellular 31 concentrations within a narrow range of concentration. 32

The mevalonate pathway is comprised of two genetic synthesis cascades which 33 react with intermediate substrates to form cholesterol and has been com-34 prehensively detailed by [22]. Sterol regulatory element-binding protein 2 35 (SREBP-2) co-regulates the gene transcription of 3-hydroxy-3-methylglutary 36 coenzyme A reductase (HMGCR) and squalene synthase. This regulation is 37 cholesterol dependent [13]. When cholesterol levels are high, SREBP-2 is 38 bound in a complex with cholesterol anchoring it to the cell membrane ren-39 dering SREBP-2 inactive. In low cholesterol concentrations the complex 40

 $_{\rm 41}$ $\,$ unbinds and through a complex series of translocation and proteolytic pro-

⁴² cessing steps SREBP-2 is released, relocates to the nucleus and binds to tar-

⁴³ get DNA stimulating increased transcription leading to increased production

⁴⁴ of the enzymes such as HMGCR and squalene synthase [6].

The central anabolic cascade of the pathway is initiated by the binding of 45 HMGCoA to the active site of HMGCR, which then catalyses its conversion 46 into mevalonate. Mevalonate is then converted to geranyl pyrophosphate 47 (geranyl-PP), farnesyl pyrophosphate (farnesyl-PP), squalene (via the inter-48 action between farnesyl-PP and squalene synthase), lanosterol and finally 49 after some 19 further steps [11], cholesterol. A rate limiting step in this 50 chain of biosynthesis is the reduction of HMGCoA catalysed by HMGCR 51 [13].52

The tight control of cholesterol concentration is thought possible by a number of negative feedback loops that regulate HMGCR and receptors dependent on intracellular cholesterol concentrations [14, 35]. Feedbacks from farnesyl-PP [10] and lanosterol accelerate HMGCR degradation [4], and it has been suggested that geranyl-PP plays a similar role. Cholesterol has been shown to accelerate squalene synthase degradation [10] and oxygenated derivatives of cholesterol have been identified in HMGCR degradation [9].

Many of the products formed from the mevalonate pathway are involved in 60 other cell signalling cascades. Farnesyl-PP is a major branch point in the 61 pathway which is responsible for producing six other substrates used in vital 62 cellular functions. Excessive amounts of farnesyl-PP have been suggestively 63 linked to tumours and Alzheimers disease [7, 32]. Inhibitors of the mevalonate 64 pathway are used in cardiovascular therapy (statins) and as anti-fungal agents 65 (CYP51 inhibitors) in crop protection. The extent to which altering this 66 pathway is associated with the carcinogenic and developmental effects of 67 CYP51 inhibitors has been debated [23, 26]. 68

Mathematical modelling of cholesterol biosynthesis pathways has to date fo-69 cused on specific aspects of the pathway. Kervizic and Corcos [19] developed 70 a boolean model of the pathway which focused on demonstrating the role of 71 SREBP-2 in synthesising cholesterol and the effect of statins on the process. 72 Their model showed good agreement with experimental known functioning of 73 the pathway in respect of statin applications. Watterson and colleagues [45] 74 formulated an ordinary differential equation (ODE) model of the pathway 75 to understand the effect of the immune response and statins on the overall 76

pathway. Using experimental data from macrophages, their work shows the 77 gradual reduction in pathway activity as a result of the innate immune re-78 sponse, versus the more step like change imparted by statins. A recent paper 79 by Bhattacharya et al. [2] formulated and analysed a three variable nonlin-80 ear ODE simplified model of the pathway that incorporates a description of 81 HMGCR mRNA, HMGCR protein and cholesterol biosynthesis. The syn-82 thesis of HMGCR mRNA is controlled by a negative feedback loop, whereby 83 cholesterol is able to bind to free SREBP-2. Model results and analysis 84 demonstrate the system exhibits one real stable steady-state which is mono-85 tonic, periodic or damped periodic under certain model parameterisations as 86 a result of cholesterol's negative regulation of SREBP-2. 87

In this paper we seek here to expand our knowledge of cholesterol biosynthesis 88 by deriving and solving a nonlinear ODE model of the mevalonate choles-89 terol biosynthesis pathway. Our aim is to better understand the role of the 90 overall network structure in dynamically regulating cholesterol biosynthesis, 91 in particular that of multiple synthesis pathways and feedbacks. We begin in 92 Section 2 by presenting our main model of the pathway which incorporates 93 the core regulation mechanisms and feedbacks within the signalling cascade. 94 An ODE model of the pathway is derived from first principles in Section 3, 95 which is subsequently parameterised and solved numerically in Section 4. 96 The results of a local sensitivity analysis are presented in Section 5 and the 97 role of the second rate limiting step in the pathway between farmesyl-PP 98 and squalene synthase is analysed in detail in Section 6. The effect of the 99 numerous feedbacks within the pathway are analysed in Section 9 before a 100 steady-state stability analysis of a model reduction of the full network model 101 is presented in Section 7. Negative feedbacks may lead to a network ex-102 hibiting oscillatory type behaviour and as such we examine whether such 103 solutions may be observed for certain parameterisations of the full model in 104 Section 8. We test the hypothesis that the application of CYP51 inhibitors 105 leads to increased levels of farnesyl-PP, via inhibition of cholesterol produc-106 tion following that of lanosterol, in Section 10. Our results and conclusions 107 are discussed in Section 11. 108

¹⁰⁹ 2. The Mevalonate Pathway

Given the complexity of the full pathway we consider here a reduction, incorporating the details outlined in the Introduction, which captures the core synthesis processes, feedbacks and branch points associated with cholesterol regulation as shown in Figure 1. Essentially, substrates and enzymes that form sequential linear steps in the pathway and which are not involved in feedbacks or branch points, have been omitted. This leaves three core aspects:

- the two genetic transcriptional control pathways of HMGCR and squalene synthase by SREBP-2;
- the central metabolic cascade which synthesises intermediary meval onate products and sterols with controlling steps using the enzymes
 HMGCR and squalene synthase; and
- negative feedback controls, including negative regulation of SREBP-2
 by cholesterol and the concentration dependent feedbacks from sterol
 and non-sterol products affecting the HMGCR and squalene synthase
 degradation rates.

In high cholesterol concentrations SREBP-2 is bound to a cholesterol molecule 126 anchoring it to the intracellular membrane, represented in Figure 1 by the 127 $\bar{\kappa}_3/\bar{\kappa}_{-3}$ negative feedback. Here $\bar{\kappa}_3$ represents the association reaction, whilst 128 $\bar{\kappa}_{-3}$ the disassociation reaction. In low cholesterol concentrations, SREBP-2 129 disassociates from the cholesterol molecule allowing it, via a series of inter-130 mediate steps, to produce an active transcription factor that relocates to the 131 nucleus to act upon the DNA stimulating endogenous production of HMGCR 132 and squalene synthase. This is represented in Figure 1, by the two reactions 133 $\bar{\kappa}_1/\bar{\kappa}_{-1}$, through $\bar{\mu}_1$ to $\bar{\mu}_3$ and $\bar{\kappa}_2/\bar{\kappa}_{-2}$, through $\bar{\mu}_2$ to $\bar{\mu}_4$. In the centre of the 134 pathway HMGCR binds with HMGCoA to form an intermediary complex 135 which leads to mevalonate production. This is subsequently phosphorylated 136 twice then converted to isopentenyl-PP and geranyl-PP. In Figure 1 these 137 five steps are represented as $\bar{\mu}_5$. From geranyl-PP, farnesyl-PP is produced. 138 It is at this point that squalene synthase reacts with farnesyl-PP and this 139 complex produces squalene. Squalene produces squalene-2,3-epoxide after 140 which lanosterol is formed. We represent these two steps by $\bar{\mu}_8$. There are a 141 further 19 reactions from lanosterol until cholesterol [11] which we approxi-142 mate by the parameter $\bar{\mu}_9$. This approximation allows for the simplification 143 of an otherwise already under parameterised system. 144

There are a number of feedbacks within the pathway shown in Figure 1. Goldstein and Brown [4] found that sterols caused a negative feedback on HMGCR production but hypothesised sterols were not the only inhibitors.



Figure 1: A simplified model of the mevalonate pathway. Arrows show forward reactions, circles show stimulative reactions and horizontal bars indicate inhibition. Here ϕ indicates the removal of a product from the pathway, either by degradation or use in another process. There are three main focal points to the pathway; the two genetic pathways of HMGCR and squalene synthase, the central metabolic cascade and the regulatory feedbacks (dashed lines).

Hence we have concentration dependent feedbacks from lanosterol (\bar{K}_7) and cholesterol (\bar{K}_8) that up-regulate the degradation of HMGCR. It has been suggested that geranyl-PP also up-regulates HMGCR degradation [14] (\bar{K}_6) and recent findings by Foresti et al. [10] have shown farnesyl-PP is linked to HMGCR degradation (\bar{K}_9) . Foresti et al. also show a similar concentration dependent reaction between cholesterol and the rate of squalene synthase degradation (\bar{K}_{10}) .

155 3. Mathematical model

1 _

In this section we derive a system of non-linear ODEs to describe the reaction
network detailed in Section 2 using the law of mass action. Details on the
biochemistry underlying each step within the pathway are given in Appendix
A. Applying the law of mass action to equations (A.1) - (A.6) gives

$$\frac{d\bar{g}_h}{d\bar{t}} = \bar{\kappa}_{-1}\bar{s}_{bh} - \bar{\kappa}_1\bar{s}^{x_h}\bar{g}_h,\tag{1}$$

$$\frac{d\bar{g}_{ss}}{d\bar{t}} = \bar{\kappa}_{-2}\bar{s}_{bss} - \bar{\kappa}_2\bar{s}^{x_s}\bar{g}_{ss},\tag{2}$$

$$\frac{ds}{d\bar{t}} = x_h\bar{\kappa}_{-1}\bar{s}_{bh} - x_h\bar{\kappa}_1\bar{s}^{x_h}\bar{g}_h + x_s\bar{\kappa}_{-2}\bar{s}_{bss} - x_s\bar{\kappa}_2\bar{s}^{x_s}\bar{g}_{ss} - \bar{\kappa}_3\bar{c}^{x_c}\bar{s} + \bar{\kappa}_{-3}\bar{c}_b, \quad (3)$$

$$\frac{ds_{bh}}{d\bar{t}} = -\bar{\kappa}_{-1}\bar{s}_{bh} + \bar{\kappa}_1\bar{s}^{x_h}\bar{g}_h,\tag{4}$$

$$\frac{d\bar{s}_{bss}}{d\bar{t}} = -\bar{\kappa}_{-2}\bar{s}_{bss} + \bar{\kappa}_2\bar{s}^{xs}\bar{g}_{ss},\tag{5}$$

$$\frac{d\bar{m}_h}{d\bar{t}} = \bar{\mu}_1 \bar{s}_{bh} - \bar{\delta}_1 \bar{m}_h, \tag{6}$$

$$\frac{dm_{ss}}{d\bar{t}} = \bar{\mu}_2 \bar{s}_{bss} - \bar{\delta}_2 \bar{m}_{ss},\tag{7}$$

$$\frac{an_r}{d\bar{t}} = \bar{\mu}_3 \bar{m}_h + \bar{\kappa}_{-4} \bar{h}_b - \bar{\kappa}_4 \bar{h}_r \bar{h}_c + \bar{\mu}_5 \bar{h}_b$$

$$- \bar{\delta}_3 \bar{h}_r \left(1 + \delta_{hg} \frac{\bar{g}_{pp}}{\bar{g}_{pp} + \bar{K}_6} + \delta_{hf} \frac{\bar{f}_{pp}}{\bar{f}_{pp} + \bar{K}_9} + \delta_{hl} \frac{\bar{l}}{\bar{l} + \bar{K}_7} + \delta_{hc} \frac{\bar{c}}{\bar{c} + \bar{K}_8} \right), (8)$$

$$\frac{d\bar{s}_s}{d\bar{s}_s} = \bar{\mu}_s \bar{m}_s + \bar{\mu}_s - \bar{f}_s - \bar{\mu}_s \bar{\bar{s}}_s - \bar{f}_s - \bar{\kappa}_s - \bar{\delta}_s - \bar{$$

$$\frac{ds_s}{d\bar{t}} = \bar{\mu}_4 \bar{m}_{ss} + \bar{\kappa}_{-5} \bar{f}_{bpp} - \bar{\kappa}_5 \bar{s}_s \bar{f}_{pp}^2 + \bar{\mu}_7 \bar{f}_{bpp} - \bar{\delta}_4 \bar{s}_s \left(1 + \delta_{sc} \frac{c}{\bar{c} + \bar{K}_{10}}\right), \tag{9}$$

$$\frac{dh_c}{d\bar{t}} = \bar{\kappa}_{-4}\bar{h}_b - \bar{\kappa}_4\bar{h}_r\bar{h}_c + \bar{\omega}, \qquad (10)$$

$$\frac{dh_b}{d\bar{t}} = -\bar{\kappa}_{-4}\bar{h}_b + \bar{\kappa}_4\bar{h}_r\bar{h}_c - \bar{\mu}_5\bar{h}_b - \bar{\delta}_3\bar{h}_b, \tag{11}$$

$$\frac{dg_{pp}}{d\bar{t}} = \bar{\mu}_5 \bar{h}_b - \bar{\delta}_5 \bar{g}_{pp} - \bar{\mu}_6 \bar{g}_{pp}, \qquad (12)$$

$$\frac{df_{pp}}{d\bar{t}} = \bar{\mu}_6 \bar{g}_{pp} - \bar{\delta}_6 \bar{f}_{pp} - 2\bar{\kappa}_5 \bar{s}_s \bar{f}_{pp}^2 + 2\bar{\kappa}_{-5} \bar{f}_{bpp}, \qquad (13)$$

$$\frac{df_{bpp}}{d\bar{t}} = \bar{\kappa}_5 \bar{s}_s \bar{f}_{pp}^2 - \bar{\kappa}_{-5} \bar{f}_{bpp} - \bar{\mu}_7 \bar{f}_{bpp} - \bar{\delta}_4 \bar{f}_{bpp}, \qquad (14)$$

$$\frac{d\bar{s}_q}{d\bar{t}} = \bar{\mu}_7 \bar{f}_{bpp} - \bar{\mu}_8 \bar{s}_q,$$

$$\frac{d\bar{l}}{d\bar{t}} = \bar{\mu}_8 \bar{s}_q - \bar{\delta}_7 \bar{l} - \bar{\mu}_9 \bar{l},$$
(15)
(16)

$$\frac{ll}{l\bar{t}} = \bar{\mu}_8 \bar{s}_q - \bar{\delta}_7 \bar{l} - \bar{\mu}_9 \bar{l}, \qquad (16)$$

$$\frac{d\bar{c}}{d\bar{t}} = \bar{\mu}_9 \bar{l} - \bar{\delta}_8 \bar{c} + x_c \bar{\kappa}_{-3} \bar{c}_b - x_c \bar{\kappa}_3 \bar{c}^{x_c} \bar{s}, \qquad (17)$$

$$\frac{d\bar{c}_b}{d\bar{t}} = \bar{\kappa}_3 \bar{c}^{x_c} \bar{s} - \bar{\kappa}_{-3} \bar{c}_b, \tag{18}$$

where, with square brackets denoting concentration, 161

$$\begin{split} \bar{g}_h &= [G_h], \quad \bar{g}_{ss} = [G_{ss}], \quad \bar{s} = [S], \quad \bar{s}_{bh} = [S_{bh}], \quad \bar{s}_{bss} = [S_{bss}], \\ \bar{m}_h &= [M_h], \quad \bar{m}_{ss} = [M_{ss}], \quad \bar{h}_r = [H_r], \quad \bar{s}_s = [S_s], \quad \bar{h}_c = [H_c], \\ \bar{h}_b &= [H_b], \quad \bar{g}_{pp} = [G_{pp}], \quad \bar{f}_{pp} = [F_{pp}], \quad \bar{f}_{bpp} = [F_{bpp}], \quad \bar{s}_q = [S_q], \\ \bar{l} = [L], \quad \bar{c} = [C], \quad \text{and} \quad \bar{c}_b = [C_b], \end{split}$$

and the system is closed with the initial conditions 162

$$\bar{g}_{h}(0) = \bar{g}_{h0}, \quad \bar{g}_{s}(0) = \bar{g}_{s0}, \quad \bar{s}(0) = \bar{s}_{0}, \quad \bar{s}_{bh}(0) = 0, \quad \bar{s}_{bss}(0) = 0, \\ \bar{m}_{h}(0) = \bar{m}_{h0}, \quad \bar{m}_{ss}(0) = \bar{m}_{ss0}, \quad \bar{h}_{r}(0) = \bar{h}_{r0}, \quad \bar{s}_{s}(0) = \bar{s}_{s0}, \\ \bar{h}_{c}(0) = \bar{h}_{c0}, \quad \bar{h}_{b}(0) = 0, \quad \bar{g}_{pp}(0) = 0, \quad \bar{f}_{pp}(0) = 0, \quad \bar{f}_{bpp}(0) = 0, \\ \bar{s}_{q}(0) = 0, \quad \bar{l}(0) = 0, \quad \bar{c}(0) = 0 \quad \text{and} \quad \bar{c}_{b}(0) = 0, \quad (19)$$

at $\bar{t} = 0$ 163

Many of the initial conditions are assumed equal to zero in order to under-164 stand the overall dynamical response of the system. The feedbacks acting 165

160

on HMGCR and squalene synthase degradation, equations (22) and (23) re-166 spectively, are dependent on geranyl-PP, farnesyl-PP, lanosterol and choles-167 terol concentrations. We thus assume these follow sigmoidal shape kinet-168 ics [24], where $K_{6,7,8,9,10}$ are the respective Michaelis-Menten constants and 169 $\delta_{hq}, \delta_{hf}, \delta_{hl}$ and δ_{hc} , are dimensionless weighting constants representing the 170 additional effect of geranyl-PP, farnesyl-PP, lanosterol and cholesterol to that 171 of the natural rate of HMGCR degradation, respectively, and δ_{sc} is that of a 172 similar effect of cholesterol on the natural decay rate of squalene synthase. 173

By invoking conservation of certain entities within the pathway and employing quasi-equilibrium approximations (see Appendix B) equations (1) to (17) are reduced to

$$\frac{d\bar{m}_h}{d\bar{t}} = \frac{\bar{\mu}_1}{1 + \left(\frac{\bar{K}_1(1 + (\frac{\bar{c}}{\bar{K}_3})^{x_c})}{\bar{s}_0}\right)^{x_h}} - \bar{\delta}_1 \bar{m}_h, \tag{20}$$

$$\frac{d\bar{m}_{ss}}{d\bar{t}} = \frac{\bar{\mu}_2}{1 + \left(\frac{\bar{K}_2(1 + (\frac{\bar{c}}{K_3})^{x_c})}{\bar{s}_0}\right)^{x_s}} - \bar{\delta}_2 \bar{m}_{ss}, \tag{21}$$

$$\frac{dh_{r}}{d\bar{t}} = \bar{\mu}_{3}\bar{m}_{h} + \bar{\kappa}_{-4}\bar{h}_{b} - \bar{\kappa}_{4}\bar{h}_{r}\bar{h}_{c} + \bar{\mu}_{5}\bar{h}_{b} \\
- \bar{\delta}_{3}\bar{h}_{r} \left(1 + \delta_{hg}\frac{\bar{g}_{pp}}{\bar{g}_{pp} + \bar{K}_{6}} + \delta_{hf}\frac{\bar{f}_{pp}}{\bar{f}_{pp} + \bar{K}_{9}} + \delta_{hl}\frac{\bar{l}}{\bar{l} + \bar{K}_{7}} + \delta_{hc}\frac{\bar{c}}{\bar{c} + \bar{K}_{8}}\right),$$
(22)

$$\frac{d\bar{s}_s}{d\bar{t}} = \bar{\mu}_4 \bar{m}_{ss} + \bar{\kappa}_{-5} \bar{f}_{bpp} - \bar{\kappa}_5 \bar{s}_s \bar{f}_{pp}^2 + \bar{\mu}_7 \bar{f}_{bpp} - \bar{\delta}_4 \bar{s}_s \left(1 + \delta_{sc} \frac{\bar{c}}{\bar{c} + \bar{K}_{10}}\right), \quad (23)$$

$$\frac{d\bar{h}_c}{d\bar{t}} = \bar{\kappa}_{-4}\bar{h}_b - \bar{\kappa}_4\bar{h}_r\bar{h}_c + \bar{\omega}, \qquad (24)$$

$$\frac{dh_b}{d\bar{t}} = -\bar{\kappa}_{-4}\bar{h}_b + \bar{\kappa}_4\bar{h}_r\bar{h}_c - \bar{\mu}_5\bar{h}_b - \bar{\delta}_3\bar{h}_b, \qquad (25)$$

$$\frac{d\bar{g}_{pp}}{d\bar{t}} = \bar{\mu}_5 \bar{h}_b - \bar{\delta}_5 \bar{g}_{pp} - \bar{\mu}_6 \bar{g}_{pp}, \qquad (26)$$

$$\frac{dJ_{pp}}{d\bar{t}} = \bar{\mu}_6 \bar{g}_{pp} - \bar{\delta}_6 \bar{f}_{pp} - 2\bar{\kappa}_5 \bar{s}_s \bar{f}_{pp}^2 + 2\bar{\kappa}_{-5} \bar{f}_{bpp}, \qquad (27)$$

$$\frac{df_{bpp}}{d\bar{t}} = \bar{\kappa}_5 \bar{s}_s \bar{f}_{pp}^2 - \bar{\kappa}_{-5} \bar{f}_{bpp} - \bar{\mu}_7 \bar{f}_{bpp} - \bar{\delta}_4 \bar{f}_{bpp}, \qquad (28)$$

177

$$\frac{d\bar{s}_q}{d\bar{t}} = \bar{\mu}_7 \bar{f}_{bpp} - \bar{\mu}_8 \bar{s}_q, \tag{29}$$

$$\frac{dl}{d\bar{t}} = \bar{\mu}_8 \bar{s}_q - \bar{\delta}_7 \bar{l} - \bar{\mu}_9 \bar{l}, \qquad (30)$$

$$\frac{d\bar{c}}{d\bar{t}} = \frac{\bar{\mu}_9 l - \delta_8 \bar{c}}{1 - x_c (\bar{s}' + x_h \bar{s}'_{bh} + x_s \bar{s}'_{bss})},\tag{31}$$

where $\bar{s}\prime_{bss}$, $\bar{s}\prime_{bh}$ and $\bar{s}\prime$ are given by equations (B.9), (B.10) and (B.8) respectively, and \prime indicates differentiation with respect to \bar{c} . The initial conditions are given by

$$\bar{m}_{h}(0) = \bar{m}_{h0}, \quad \bar{m}_{ss}(0) = \bar{m}_{ss0}, \quad \bar{h}_{r}(0) = \bar{h}_{r0}, \quad \bar{s}_{s}(0) = \bar{s}_{s0}, \\ \bar{h}_{c}(0) = \bar{h}_{c0}, \quad \bar{h}_{b}(0) = 0, \quad \bar{g}_{pp}(0) = 0, \quad \bar{f}_{pp}(0) = 0, \quad \bar{f}_{bpp}(0) = 0, \\ \bar{s}_{q}(0) = 0, \quad \bar{l}(0) = 0 \quad \text{and} \quad \bar{c}(0) = 0.$$
(32)

181 3.1. Non-dimensionalisation

Equations (20) to (32) are non-dimensionalised according to the following rescalings

$$\bar{t} = \frac{t}{\delta_7}, \quad \bar{m}_h = \bar{m}_{h0} m_h, \quad \bar{m}_{ss} = \bar{m}_{h0} m_{ss}, \quad \bar{h}_r = \bar{s}_{sT} h_r,$$
$$\bar{S}_s = \bar{s}_{sT} s_s, \quad \bar{h}_c = \bar{h}_{cT} h_c, \quad \bar{h}_b = \bar{h}_{cT} h_b, \quad \bar{g}_{pp} = \bar{h}_{cT} g_{pp},$$
$$\bar{f}_{pp} = \bar{h}_{cT} f_{pp}, \quad \bar{f}_{bpp} = \bar{h}_{cT} f_{bpp}, \quad \bar{s}_q = \bar{h}_{cT} s_q, \quad \bar{l} = \bar{h}_{cT} l, \quad \bar{c} = \bar{h}_{cT} c, \quad (33)$$

where \overline{s}_{sT} and \overline{h}_{cT} are the experimentally determined total concentrations of squalene synthese and HMG-CoA in a resting hepatocyte cell [5]. Substitut $_{186}$ ing these rescalings into equations (20) through (32), we obtain

$$\frac{dm_h}{dt} = \frac{\mu_1}{1 + \left(\kappa_1 (1 + (\frac{c}{\kappa_3})^{x_c})\right)^{x_h}} - \delta_1 m_h, \tag{34}$$

$$\frac{dm_{ss}}{dt} = \frac{\mu_2}{1 + \left(\kappa_2 (1 + (\frac{c}{\kappa_3})^{x_c})\right)^{x_s}} - \delta_2 m_{ss}, \tag{35}$$

$$\frac{dh_r}{dt} = \mu_3 m_h + \kappa_{-4} \alpha h_b - \kappa_4 \alpha h_r h_c + \mu_5 \alpha h_b
-\delta_3 h_r \left(1 + \delta_{hg} \frac{g_{pp}}{g_{pp} + K_6} + \delta_{hf} \frac{f_{pp}}{f_{pp} + K_9} + \delta_{hl} \frac{l}{l + K_7} + \delta_{hc} \frac{c}{c + K_8} \right),$$
(36)

$$\frac{ds_s}{dt} = \mu_4 m_{ss} + \kappa_{-5} \alpha f_{bpp} - \kappa_5 \alpha s_s f_{pp}^2 + \mu_7 \alpha f_{bpp} - \delta_4 s_s \left(1 + \delta_{sc} \frac{c}{c + K_{10}}\right),$$
(37)

$$\frac{dh_c}{dt} = \kappa_{-4}h_b - \kappa_4h_rh_c + \omega, \qquad (38)$$

$$\frac{dh_b}{dt} = -\kappa_{-4}h_b + \kappa_4h_rh_c - \mu_5h_b - \delta_3h_b, \qquad (39)$$

$$\frac{dt}{dg_{pp}} = \mu_5 h_b - \delta_5 g_{pp} - \mu_6 g_{pp}, \tag{40}$$

$$\frac{dg_{pp}}{dt} = \mu_6 g_{pp} - \delta_6 f_{pp} - 2\kappa_5 s_s f_{pp}^2 + 2\kappa_{-5} f_{bpp}, \qquad (41)$$

$$\frac{df_{bpp}}{dt} = \kappa_5 s_s f_{bpp}^2 - \kappa_{-5} f_{bpp} - \mu_7 f_{bpp} - \delta_4 f_{bpp}, \qquad (42)$$

$$\frac{ds_q}{dt} = \mu_7 f_{bpp} - \mu_8 s_q, \tag{43}$$

$$\frac{dl}{dt} = \mu_8 s_q - \delta_7 l - \mu_9 l, \tag{44}$$

$$\frac{dl}{dt} = \mu_8 s_q - \delta_7 l - \mu_9 l,$$
(44)
$$\frac{dc}{dt} = \frac{\mu_9 l - \delta_8 c}{1 - x_c (s_0 s' + x_h g_{h0} s'_{bh} + x_s g_{ss0} s'_{bss})},$$
(45)

with the non-dimensional initial conditions, at t = 0, given by

$$m_h(0) = 1, \quad m_{ss}(0) = 1, \quad h_r(0) = 0, \quad s_s(0) = 0, \quad h_c(0) = 0,$$

$$h_b(0) = 0, \quad g_{pp}(0) = 0, \quad f_{pp}(0) = 0, \quad f_{bpp}(0) = 0, \quad s_q(0) = 0,$$

$$l(0) = 0 \quad \text{and} \quad c(0) = 0, \quad (46)$$

¹⁸⁹ and the non-dimensional parameters summarised in Table 2.

¹⁹⁰ 3.2. Model parameterisation

Wherever possible data from human liver (hepatocyte G2; HepG2) cells 191 was used to inform our parameter values. Where values have been unavail-192 able from HepG2 cells, other sources have included human liver microsomes 193 (pieces of the endoplasmic reticulum used in some experimental work) or Chi-194 nese hamster ovary cells. Details regarding the estimation of all parameter 195 values is provided in Appendix C, whilst Table 1 summarises each dimen-196 sional parameter, their value and source. Non-dimensional parameters are 197 stated in Table 2. 198

In cases where no information was available, approximations were first made 199 based on similar occuring reactions and processes, e.g. rates of mRNA 200 degradation, as detailed in Appendix C. For instance, rates calculated from 201 Bhattacharya et al. [2] regarding HMGCR and cholesterol synthesis, specif-202 ically binding affinities and degradation rates relating to HMGCR mRNA, 203 HMGCR and cholesterol, were used to initially inform rates corresponding 204 to squalene synthesis and degradation as well as (non)sterol pro-205 duction rates. Using Matlab [21] the model was then simulated numerically 206 (using the ode15s solver) and analysed via a local sensitivity analysis (coded 207 directly into Matlab). The sensitivity analysis was used to ascertain the 208 importance of the unknown assumed parameter values in affecting the to-209 tal cholesterol concentration in an heptaocyte. Based on the findings of this 210 analysis, parameter values were then adjusted accordingly (as detailed in Ap-211 pendix C) to ensure the model reproduced previously determined cholesterol 212 concentrations [2]. 213

²¹⁴ In the absence of any available data in other cell systems with which to ²¹⁵ compare any determined values, the additional effects of farnesyl-PP, geranyl-²¹⁶ PP, lanosterol and cholesterol on HMGCR degradation and cholesterol on that of squalene synthase degradation (δ_{hg} , δ_{hf} , δ_{hl} , δ_{hc} , δ_{sc}) were set equal to unity.

It is important to note that the utilisation of cholesterol and farnesyl-PP can
vary depending on other intracellular processes. To simplify our model, we
have assumed a constant value of cholesterol and farnesyl-PP degradation to

include cellular utilisation, based on the work by Bhattacharya et al. [2].

D	Description	V-l	II	Defense
Param.	Description	Value	Units	Reference
\bar{m}_{h0}	Initial HMGCR mRNA	3.0×10^{9}	molec./ml	[30]
	concentration.			
\bar{m}_{ss0}	Initial SqS mRNA	$3.0 imes 10^9$	molec./ml	[30]
	concentration.			
\bar{s}_{sT}	Total SqS synthase	7.59×10^{14}	molec./ml	[5]
	concentration.			
\bar{h}_{cT}	Total HMGCoA	$1.98 imes 10^{15}$	molec./ml	[33, 38]
	concentration.			
\overline{s}_0	Total SREBP-2	8.21×10^{16}	molec./ml	[31, 2]
	concentration.		,	
\bar{g}_{h0}	HMGCR gene	2.11×10^9	molec./ml	[41]/This study.
0	concentration.		,	
\bar{q}_{ss0}	SqS gene	2.11×10^{9}	molec./ml	This study.
J330	concentration.	-		
$\bar{\mu}_1^*$	HMGCR transcription.	5.17×10^5	molec.	[8, 12]
, 1 *	CaC transcription	4.65×10^5	mi.s molec.	[0 27]
μ_2	SqS transcription.	4.00×10^{-5}	ml.s	[8, 37]
$ar{\mu}_3$	HMGCR translation	3.32×10^{-2}	1/s	[39, 17]
$ar{\mu}_4$	SqS translation.	1.91×10^{-2}	1/s	[39, 36]
$ar{\mu}_5$	Geranyl-PP formation.	$4.33 imes 10^{-2}$	1/s	[15, 33, 43]
$ar{\mu}_6$	Farnesyl-PP formation.	$4.33 imes10^{-2}$	1/s	[15, 33, 47]
$\bar{\mu}_7$	SqS formation.	2.17×10^{-1}	1/s	This study.
$\bar{\mu}_8$	Lanosterol formation.	4.33×10^{-2}	1/s	[15, 33, 47]
$\bar{\mu}_9$	Cholesterol formation.	4.33×10^{-2}	1/s	[15, 33, 47]
\overline{K}_1	SREBP-2-HMGCR gene	8.21×10^{12}	molec./ml	[29]/This study.
-	binding affinity.		,	
\bar{K}_2	SREBP-2-SaS gene	8.21×10^{12}	molec./ml	[29]/This study.
2	binding affinity.			
\bar{K}_2	Cholesterol-SREBP-2	1.49×10^{16}	molec /ml	[46]/This study
110	disassociation constant	1.10 / 10		
Ē.	HMGCB-HMGCoA	1.39×10^{-16}	l	This study
n_4	aga sistion	1.00×10	molec.s	1 mo soury.
	association.			

Table 1: Dimensional parameters. Here "Param." denotes parameter, "molec
" molecules, "SqS" squalene synthase.

	Table 1	continued		
$\bar{\kappa}_{-4}$	HMGCR-HMGCoA	1.75×10^{-7}	1/s	This study
	disassociation.		• 2	
$\bar{\kappa}_5$	SqS - Farnesyl-PP	1.76×10^{-30}	$\frac{\text{ml}^2}{\text{molec}^2.\text{s}}$	This study
	association.			
$\bar{\kappa}_{-5}$	SqS - Farnesyl-PP	1.75×10^{-5}	1/s	This study.
_	disassociation.			
K_6	Michaelis-Menten constant	5.00×10^{9}	molec./ml	This study.
	for geranyl-PP/HMGCR			
_	degradation.			
\overline{K}_7	Michaelis-Menten constant	5.00×10^{12}	molec./ml	This study.
	for lanosterol/HMGCR			
_	degradation.			
K_8	Michaelis-Menten constant	5.00×10^{17}	molec./ml	This study.
	for cholesterol/HMGCR			
_	degradation.			
\overline{K}_9	Michaelis-Menten constant	5.00×10^{11}	molec./ml	This study.
	for farnesyl-PP/HMGCR			
_	degradation.			
\overline{K}_{10}	Michaelis-Menten constant	5.00×10^{17}	molec./ml	This study.
	for cholesterol/SqS			
_	degradation.			
δ_1	HMGCR mRNA	4.48×10^{-5}	1/s	[3]
_	degradation.	_		
δ_2	SqS mRNA	4.48×10^{-5}	1/s	This study.
_	degradation.	_		
δ_3	HMGCR degradation.	6.42×10^{-5}	1/s	[44]
$\frac{\delta_4}{2}$	SqS degradation.	6.42×10^{-5}	1/s	This study.
$\frac{\delta_5}{2}$	Geranyl-PP degradation.	1.20×10^{-4}	1/s	This study.
$\frac{\delta_6}{2}$	Farnesyl-PP degradation.	1.20×10^{-4}	1/s	This study.
$\frac{\delta_7}{2}$	Lanosterol degradation.	1.20×10^{-4}	1/s	This study.
δ_8	Cholesterol degradation.	1.20×10^{-4}	1/s	[2]
δ_{hg}	Additional effect of geranyl-PP	1	-	This study
	on HMGCR degradation.		•	•
δ_{hf}	Additional effect of farmesyl-PP	1	-	This study
_	on HMGCR degradation.		•	
δ_{hl}	Additional effect of lanosterol	1	-	This study
	on HMGCR degradation.			
δ_{hc}	Additional effect of cholesterol	1	-	This study
-	on HMGCR degradation.	_		
δ_{sc}	Additional effect of cholesterol	1	-	This study
	on SqS degradation.	0.0011	• • •	
$\bar{\omega}$	HMGCoA production.	3.90×10^{11}	molec./ml	This study.
x_h	Binding sites on HMGCR	3	-	[[28]

Table 1 – continued

Table 1 – continued

	gene for SREBP-2.			
x_s	Binding sites on SqS	1	-	This study.
	gene for SREBP-2.			
x_c	Molecules of cholesterol	4	-	[46, 16]
	to inactivate SREBP-2.			

Table 2:	Table of non-dimensional	parameters,	their	relation	to	dimensional	ones	and
value.								

Parameter	Description	Definition	Value
s_0	Ratio of SREBP-2 to HMGCoA	\bar{s}_0/\bar{h}_{cT}	41.46
g_{h0}	Ratio of HMGCR gene to SREBP-2	\bar{g}_{h0}/\bar{s}_0	2.57×10^{-8}
g_{ss0}	Ratio of SqS gene to SREBP-2	\bar{g}_{ss0}/\bar{s}_0	2.57×10^{-8}
μ_1	HMGCR mRNA transcription.	$rac{ar{\mu}_1^-}{ar{\delta}_7ar{m}_{h0}}$	1.44
μ_2	SqS mRNA transcription.	$\frac{\bar{\mu}_2^*}{\bar{\delta}_7 \bar{m}_{h0}}$	1.29
μ_3	HMGCR translation.	$\frac{\bar{\mu}_3 \bar{m}_{h0}}{\bar{\delta}_7 \bar{s}_s T}$	1.10×10^{-3}
μ_4	SqS translation.	$\frac{\bar{\mu}_4 \bar{m}_{h0}}{\bar{\delta}_7 \bar{s}_{\circ} T}$	6.29×10^{-4}
μ_5	Geranyl-PP production.	$\frac{\overline{\mu}_{5}}{\overline{\delta}_{7}}$	3.61×10^2
μ_6	Farnesyl-PP production.	$\frac{\overline{\mu}_{6}}{\overline{\delta}_{7}}$	$3.61 imes 10^2$
μ_7	SqS production.	$\frac{\frac{\overline{\mu}_{7}}{\overline{\lambda}_{7}}}{\overline{\lambda}_{7}}$	1.80×10^3
μ_8	Lanosterol production.	$\frac{\frac{\overline{\mu}}{2}}{\overline{\delta}_{\pi}}$	3.61×10^2
μ_9	Cholesterol production.	$\frac{\overline{\mu}_{9}}{\overline{\delta}_{7}}$	3.61×10^2
κ_1	SREBP-2-HMGCR gene binding affinity.	$\frac{\bar{K}_1}{\bar{s}_0}$	1×10^{-4}
κ_2	SREBP-2-SqS gene	$\frac{\overline{\tilde{K}}_2}{\overline{\tilde{s}}_2}$	1×10^{-4}
	binding affinity.	50	
κ_3	Cholesterol-SREBP-2 dissociation	$\frac{\bar{K}_3}{\bar{h}_{oT}}$	7.5
	constant.	1021	
κ_4	HMGCR-HMGCoA association.	$\frac{\bar{\kappa}_4 \bar{s}_{sT}}{\bar{\delta}_7}$	8.83×10^2
κ_{-4}	HMGCR-HMGCoA disassociation.	$\frac{\overline{\kappa}_{-4}}{\overline{\delta}_{7}}$	1.46×10^{-3}
κ_5	SqS-farnesyl-PP	$\frac{\bar{\kappa}_5 \bar{h}_{cT} \bar{s}_{sT}}{\bar{\delta}_7}$	2.20×10^4
	association.		
κ_{-5}	SqS-farnesyl-PP	$\frac{\overline{\kappa}_{-5}}{\overline{\delta}_{7}}$	1.46×10^{-1}
	disassociation.		
K_6	Michaelis-Menten constant	$\frac{K_6}{\bar{h}_{cT}}$	2.53×10^{-6}
	for geranyl-PP/HMGCR degradation.		
K_7	Michaelis-Menten constant	$\frac{\overline{K}_7}{\overline{h}_{oT}}$	2.53×10^{-3}
	for lanosterol/HMGCR degradation.		
K_8	Michaelis-Menten constant	$\frac{\bar{K}_8}{\bar{h}_c T}$	2.53×10^2
	for cholesterol/HMGCR degradation.	10C1	
K_9	Michaelis-Menten constant	$\frac{\bar{K}_9}{\bar{h}_{cT}}$	2.53×10^{-4}

Table 2 – continued					
	for farnesyl-PP/HMGCR degradation.				
K_{10}	Michaelis-Menten constant	$\frac{K_{10}}{\bar{h}_{-T}}$	$2.53 imes 10^2$		
	for cholesterol/SqS degradation.	<i>nc1</i>			
δ_1	HMGCR mRNA degradation.	$\frac{\overline{\delta}_1}{\overline{\delta}_7}$	$3.73 imes 10^{-1}$		
δ_2	SqS mRNA degradation.	$\frac{\overline{\delta}_2}{\overline{\delta}_7}$	3.73×10^{-1}		
δ_3	HMGCR degradation.	$\frac{\overline{\delta}_3}{\overline{\delta}_7}$	$5.35 imes 10^{-1}$		
δ_4	SqS degradation.	$\frac{\overline{\delta_4}}{\overline{\delta_7}}$	$5.35 imes 10^{-1}$		
δ_5	Geranyl-PP degradation.	$\frac{\overline{\delta}_{5}}{\overline{\delta}_{7}}$	1		
δ_6	Farnesyl-PP degradation.	$\frac{\overline{\delta}_{6}}{\overline{\delta}_{7}}$	1		
δ_7	Lanosterol degradation.	$\frac{\overline{\delta}_7}{\overline{\delta}_7}$	1		
δ_8	Cholesterol degradation.	$\frac{\overline{\delta}_8}{\overline{\delta}_7}$	1		
ω	HMGCoA production.	$\frac{\overline{\vec{\omega}}}{\overline{\delta_7 h_{a0}}}$	0.82		
α	Ratio of total HMGCoA to SqS.	$\frac{\underline{h}_{cT}}{\overline{s}_{sT}}$	2.61		

4. Analysis of numerical results

In this section we present numerical solutions to equations (34) to (46), parameterised by Table 2, obtained using the MATLAB stiff differential equation solver ode15s [21]. Results are shown in Figure 2. Time has been redimensionalised on the x-axis and simulations run until the system reaches steady-state.

Figure 2 shows the initial increase of HMGCR and squalene synthese mRNA; 229 a result of no cholesterol being initially present in the system. HMGCR and 230 squalene synthase mRNA transcription subsequently leads to their transla-231 tion into their respective proteins. As HMGCR increases it binds to HMG-232 CoA leading to a subsequent decrease in its levels. This substrate-enzyme 233 reaction leads to increases in geranyl-PP, farnesyl-PP, bound farnesyl-PP 234 with squalene synthase, squalene, lanosterol and finally cholesterol. The ob-235 served decrease in each entity within the network at approximately 20 hours 236 is the result of global and local feedbacks within the system. Firstly, the in-237 crease in cholesterol leads, via the negative feedback between cholesterol and 238 SREBP-2 transcription of HMGCR mRNA and squalene synthese mRNA, 239 to a decrease in the concentration of HMGCR and squalene synthese, re-240 spectively. This globally controlled feedback reduction in the two enzymes 241 subsequently means less of the central cascade products are now being synthe-242 sised. This feedback is explored in more detail in Section 7.1. Simultaneously, 243



Figure 2: Numerical solutions to equations (34) to (46) with parameter values detailed in Table 2. Solutions show the response of HMGCR and squalene synthase mRNA to initial zero cholesterol concentrations, the subsequent increase in HMGCR and squalene synthase which allows the synthesis of cascade products geranyl-PP, farnesyl-PP, squalene, lanosterol and finally cholesterol.

and more locally, negative feedbacks from geranyl-PP, farnesyl-PP, lanosterol
and cholesterol seek to limit the enzymatic action of HMGCR and squalene
synthase by increasing their rates of degradation. These local feedbacks are
explored in more detail in Section 9.

The subsequent decrease in cholesterol levels leads to a small increase in 248 HMGCR and squalene synthase mRNA transcription. Eventually the so-249 lutions evolve to reach a steady-state. Solutions of the model show that 250 concentrations of both farnesyl-PP and cholesterol are greater than those of 251 other cascade products; geranyl-PP, squalene and lanosterol. One reason for 252 this could be because farnesyl-PP is a major branch point in the pathway 253 and is used (as is cholesterol) in a greater number of cell processes, thus their 254 respective concentrations need to be higher. We note that HMGCoA initially 255 increases (as a result of its own synthesis) before decreasing to steady-state 256 levels due to increased HMGCR levels. 257

Direct comparison with experimental values for the concentration of each en-258 tity within the pathway is difficult given a lack of reported values in the lit-259 erature. In the case of HMGCR mRNA we can approximate this via Rudling 260 et al. [30] who states there are 30 copies of HMGCR mRNA found in each 261 human liver cell under basal conditions. This leads to a concentration of 262 3.00×10^{10} molecules/ml, for which our result of 1.13×10^{10} molecules/ml is 263 very similar. Our concentrations of HMGCR mRNA, HMGCR and choles-264 terol are also in agreement with those previously reported in Bhattacharya 265 et al. [2]. 266

²⁶⁷ 5. Model Analysis

In this and subsequent sections we undertake a comprehensive analysis of the 268 mevalonate pathway model. Given the overall network complexity and diffi-269 culty in obtaining analytical solutions to the system of governing equations 270 we begin with a sensitivity analysis in Section 5.1. Results from this high-271 light enzyme-rate rate limiting steps within the pathway which are explored 272 in more detail analytically in Section 6. We consider a simplified model of 273 the pathway, containing the key enzyme-substrate reactions and feedbacks 274 within the pathway in Section 7, in order to examine the steady-states of the 275 system and their stability. Numerical experiments are conducted in Section 8 276 to verify our findings. 277

278 5.1. Sensitivity analysis

We conducted a local sensitivity analysis, varying each parameter in turn, up 279 to 100-fold above and below the values reported in Table 2. We quantitatively 280 measured, primarily, the effect of parameter variation on the steady-state 281 cholesterol concentrations (in relation to the unperturbed system) whilst also 282 looking for significant variations in key elements of the pathway, for exam-283 ple steady-state farnesyl-PP concentrations and differences in the dynamic 284 behaviour of each model variable. Varying the model parameters up to 100-285 fold allows us to explore the robustness of the pathway to changes greater 286 than those biologically feasible thereby ensuring all possible effects have been 287 explored. 288

In what follows we present our results by discussing parameters related to specific processes within the pathway (e.g. HMGCR synthesis) wherever possible. Given their number and to ascertain their effects separately, negative feedbacks within the pathway are discussed separately in Sections 7.1 and 9. Not all parameters caused a notable change in the system; only those that did are discussed here.

The results of our local sensitivity analysis were subsequently confirmed by a metabolic control analysis in which the relationship between the system steady-states and the properties of the individual reactions was explored. The response coefficients were calculated via

$$\mathbf{R} = R_m^i = \frac{P_m}{S_i^{st}} \frac{\partial S_i^{st}}{\partial P_m},$$

where **R** is the matrix of response coefficients, P_m is each parameter value and S_i^{st} is the corresponding metabolite (mRNA/substrate/enzyme in our system) at steady-state.

298 5.1.1. HMGCoA synthesis (ω)

The HMGCoA-HMGCR reaction point in the pathway is the first rate limiting step in the cascade [34] and HMGCoA is the starting point of all the central cascade reactions. Hence decreasing HMGCoA availability 10-fold leads to an abundance of enzyme HMGCR (over 300% more) and leads to a reduction of over 90% in all cascade products except farnesyl-PP (73%).

Increasing the rate of HMGCoA synthesis 10-fold, decreases HMGCR con-304 centrations by almost 100% due to the abundance of HMGCoA, but has only 305 a moderate effect on the concentrations of cascade products (around 33%) 306 including cholesterol. In all cases farnesyl-PP is more tightly regulated, and 307 exhibits a smaller percentage change, than the rest of the cascade products. 308 Thus the farnesyl-PP squalene synthese substrate-enzyme reaction appears 309 to act as a second rate limiting step in the pathway, lending greater control 310 to downstream cholesterol concentrations. This is explored in further detail 311 in Section 6. 312

313 5.1.2. Genetic regulation of HMGCR (μ_1 , μ_3 , δ_1 and δ_3)

Parameter changes that induce an increase in HMGCR mRNA or HMGCR 314 did not greatly affect the pathway. This is because the substrate HMGCoA is 315 almost completely utilised and thus cholesterol increases are limited in spite 316 of the amount of HMGCR being produced i.e. the binding of HMGCoA 317 and HMGCR has reached its upper limit. This combined with the results of 318 altering the rate of HMGCoA synthesis ω , show there is a careful balance of 319 both enzyme HMGCR and substrate HMGCoA in order for cholesterol to be 320 produced. If there is an abundance of either enzyme or substrate, the reaction 321 will be limited by the lower of the two concentrations without a significant 322 effect on cholesterol concentrations. However, biologically, we would always 323 expect the concentration of enzyme to be less than the concentration of 324 substrate. 325

On the other hand, decreasing the rates of transcription and translation (μ_1 , 326 μ_3) or increasing the rates of HMGCR mRNA and HMGCR degradation (δ_1 327 and δ_3) has a significant effect on cholesterol concentrations, as well as de-328 creasing all the other cascade products. For example, decreasing the value of 329 μ_1 or μ_3 by even one order of magnitude causes an 88% decrease in cholesterol 330 levels. Increasing the value of δ_1 or δ_3 by one order of magnitude has a simi-331 lar effect. Concentrations of HMGCR are, unsurprisingly, decreased leading 332 to an accumulation of HMGCoA. Products of the central cascade are all 333 decreased by around 88% (farnesyl-PP 68%). The reduction of cholesterol 334 upregulates squalene synthase via the local squalene synthase degradation 335 feedback shown in Figure 1. 336

337 5.2. Genetic regulation of squalene synthase $(\mu_2, \mu_4, \delta_2 \text{ and } \delta_4)$

Parameter changes that cause an increase in squalene synthase mRNA or 338 squalene synthase do not greatly affect the pathway. An abundance in squa-339 lene synthase leads to a significant decrease in farnesyl-PP, but the increase 340 in cholesterol concentrations (as well as those of squalene and lanosterol) is 341 only around 7%. Increasing the amount of squalene synthase does have a 342 greater effect on cholesterol concentrations than increasing the amount of 343 HMGCR, however we again see the balanace of enzyme and substrate limit-344 ing the reaction. 345

Parameter changes that cause a decrease in squalene synthase mRNA or 346 squalene synthase have less of an effect on concentrations of cholesterol than 347 a decrease in HMGCR. For example, decreasing the value of transcription of 348 squalene synthase mRNA (μ_2) or translation of squalene (μ_4), by one order of 349 magnitude causes a 39% decrease in cholesterol levels. Increasing the value of 350 δ_2 (the degradation rate of squalene synthase mRNA) or δ_4 (the degradation 351 rate of squalene synthase) by one order of magnitude has the same effect. In 352 each case concentrations of squalene synthase are, unsurprisingly, decreased 353 which leads to an accumulation of farnesyl-PP. Products downstream of the 354 farnesyl-PP-squalene synthase reaction (bound farnesyl-PP, squalene, lanos-355 terol and cholesterol) are all decreased by around 39%, another indicator of 356 a limiting step at this point in the pathway. This decline in cholesterol and 357 other cascade product concentrations slightly reduces HMGCR degradation 358 (2% change) as expected. We can demonstrate the effect of the HMGCR 359 degradation feedbacks by comparing the concentrations between one and 360 two orders of magnitude change in δ_2 and δ_4 , where cholesterol and lanos-361 terol concentrations decrease by 92.6%, HMGCR concentrations increase by 362 8%. 363

5.2.1. Association and disassociation of HMGCR for HMGCoA and farnesyl-PP for squalene (κ_4 , κ_{-4} , κ_5 and κ_{-5})

Altering the association rates of each of these enzyme and substrate reactions has a small effect on cholesterol levels and downstream cascade products. We found that decreasing the rate of binding (κ_5) in the squalene synthasefarnesyl-PP reaction, has a greater effect on cholesterol and downstream cascade product levels than decreasing the binding rate (κ_4) in the HMGCR-HMGCoA reaction, again indicating the importance of the squalene synthasefarnesyl-PP rate limiting step. Altering the disassociation rates (κ_{-4} and κ_{-5}) of each reaction has no effect on cholesterol levels or indeed the rest of the system.

375 5.2.2. Production of geranyl-PP and squalene (μ_5 and μ_7)

Decreasing the rate at which either of the enzyme-substrate complexes are 376 converted to a product decreases the concentrations of the respective down-377 stream products. Specifically decreasing the rate of squalene production, has 378 a lesser effect on products downstream of the reaction than decreasing the 379 rate at which geranyl-PP is produced. Decreasing either of these rates leads 380 to an increase in both substrate concentrations but, counter-intuitively, de-381 creases the concentration of both enzymes. This happens for two reasons; 382 firstly the enzymes are held in their bound rather than free forms (shown by 383 an increase in bound substrate concentrations). Secondly, increases in each 384 substrate concentration ensures that any enzyme synthesised or returned 385 from disassociation with the enzyme-substrate complex is quickly bound by 386 the excess substrate. Increasing the rate of complex to product conversion 387 $(\mu_5 \text{ and } \mu_7)$ has very little effect on downstream cascade products, given they 388 are limited by the amount of available substrate (HMGCoA and farnesyl-PP, 389 respectively). 390

³⁹¹ 5.2.3. Production of farnesyl-PP and lanosterol (μ_6 and μ_8)

Increasing the production rate of farnesyl-PP and lanosterol has very little effect on the pathway and cholesterol levels. Interestingly, decreasing the production rate of farnesyl-PP has a greater effect on the central cascade products than decreasing the production of lanosterol. Decreasing μ_6 100fold reduces cholesterol concentrations by 22%, reducing the degradation of HMGCR and squalene synthase, which increase by 1.5% and 3.4% respectively.

5.2.4. Production of cholesterol (μ_9)

Increasing the rate of production of cholesterol does not greatly affect cholesterol concentrations, however decreasing μ_9 has a small to moderate effect on cholesterol levels. However, the changes in lanosterol concentrations as a result, have the greatest effect on HMGCR concentrations via the local degradation feedbacks, in comparison to parameter changes that induce an
increase or reduction of geranyl-PP or farnesyl-PP - the other degradation
feedbacks on HMGCR.

407 5.2.5. Degradation of farnesyl-PP (δ_6)

Decreasing the degradation rate of farnesyl-PP slightly increases the steady-408 state concentration of cholesterol and other downstream cascade products 409 (within 10%). As expected this negatively effects both HMGCR and squalene 410 synthase via the degradation feedbacks by a moderate amount in order to 411 limit the increase in farnesyl-PP and cholesterol. However, increasing the 412 degradation rate of farnesyl-PP by just one order of magnitude impacts the 413 downstream cascade significantly, decreasing the concentrations of squalene, 414 lanosterol and cholesterol by 52.4% (33.6% for farnesyl-PP). The decrease in 415 cholesterol subsequently up-regulates HMGCR and squalene synthase levels. 416 Interestingly, squalene synthase is increased slightly more than HMGCR. 417 This could be to counteract the loss of farnesyl-PP through degradation, to 418 ensure cholesterol concentrations are maintained. 419

⁴²⁰ 5.2.6. Degradation of geranyl-PP and lanosterol (δ_5 and δ_7)

Altering the degradation rates of geranyl-PP and lanosterol have very little 421 effect on the pathway or steady-state cholesterol levels. Increasing degra-422 dation of geranyl-PP by 100 fold moderately reduces the concentrations of 423 the central cacade and slightly upregulates squalene synthase and HMGCR. 424 Squalene synthase more so. Increasing the degradation rate of lanosterol by 425 100-fold also reduces the concentrations of lanosterol and cholesterol by ap-426 proximately the same amount, however, HMGCR is upregulated more than 427 squalene synthase. This is a result of the change in central cascade products 428 and the role of the Michaelis-Menten responses affecting the feedbacks to 429 HMGCR and squalene synthase, respectively. 430

431 5.2.7. Cholesterol degradation (δ_8)

Varying the rate of cholesterol degradation greatly effects cholesterol concentrations. As expected the increase in cholesterol concentrations downregulates HMGCR and squalene synthase via the local degradation feedbacks,

however only by around 1%. Similarly for decreased cholesterol concentrations, HMGCR and squalene synthse are upregulated by around 0.1%.

437 5.2.8. Genetic binding affinities and stoichiometric coefficients (κ_1 , κ_2 , κ_3 , 438 x_c , x_h and x_s)

Binding affinities and stoichiometric coefficients involved with the genetic 439 regulation of HMGCR and squalene synthase have very little effect on the 440 system. Interestingly, reducing parameters involved in genetic regulation of 441 HMGCR has a greater effect on the system than those in regulating squalene 442 synthase, however these changes would indicate a fraction of a binding site 443 which is biologically infeasible. Furthermore, decreasing the value of κ_3 , the 444 regulation of HMGCR and squalene synthase by cholesterol has the effect 445 of decreasing cholesterol concentrations, significantly for a 100-fold decrease, 446 whilst slightly upregulating HMGCR and squalene synthase. 447

448 5.2.9. Sensitivity analysis summary

Local sensitivity analysis has highlighted that a decrease in HMGCR (the 449 first rate limiting step in the pathway), caused by parameters linked with 450 its genetic regulation, significantly decreases steady-state cholesterol concen-451 trations. However, increases in products linked with genetic regulation of 452 HMGCR do not have a significant impact on steady-state cholesterol con-453 centrations, due to the occurrence of the second rate limiting step between 454 squalene synthase and farnesyl-PP. The effect of decreasing products linked 455 with genetic regulation of squalene synthase is not as significant as those 456 linked with regulation of HMGCR. 457

An increase in products prior to the reaction of farnesyl-PP with squalene 458 synthase rarely causes a significant change in cholesterol levels (the excep-459 tion being a decrease in μ_5 reducing cholesterol concentrations significantly), 460 whilst the degradation of farnesyl-PP has a high effect on downstream prod-461 uct concentrations. We found that, with the exception of decreasing μ_5 , the 462 rates of geranyl-PP and squalene formation, from the two enzyme-substrate 463 reactions within the pathway, have a moderate effect on limiting downstream 464 products formed in the pathway. In contrast, altering the rates of geranyl-465 PP and lanosterol degradation have little impact on the pathway. Cellular 466 cholesterol concentrations are very sensitive to changes in the rate of choles-467

terol esterification (degradation) without much interruption to the rest of the pathway.

Our results, as summarised in Table 3, demonstrate that the two rate limiting 470 steps of HMGCR and HMGCoA and farnesyl-PP and squalene synthase, 471 coupled with the negative feedback between cholesterol and SREBP-2, act 472 as core regulators of products within the central cascade. The HMGCoA 473 and HMGCR rate limiting step is aimed at controlling production of central 474 cascade substrates, whilst that of farnesyl and squalene synthase appears 475 two-fold; it acts to control the levels of lanosterol and ultimately cholesterol 476 produced, but also regulate those of farnesyl-PP, given its role in other cell 477 signalling pathways. Whilst the enzyme rate limiting step of HMGCR and 478 HMGCoA follows one-to-one stoichometry, this differs for squalene synthase 479 and farnesyl-PP; two molecules of farnesyl-PP reversibly bind to squalene 480 synthase, to produce one molecule of complex bound farnesyl-PP. The effect 481 of this is investigated further in Section 6. 482

Table 3: Sensitivity analysis summary. Results here indicate up to a 10% (denoted '+' or '-'), 10-50% ('++/- -'), greater than 50% ('+++/- - -') variation or no change ('NC') in the steady-state cholesterol levels for the parameterisation detailed in Table 1 for 10-fold parameter variations.

Change Made	Parameters Involved	Effect on Cholesterol
Increased HMGCoA	ω	++
Decreased HMGCoA	ω	
Increased HMGCR	$\mu_1,\mu_3,\delta_1,\delta_3$	NC
Decreased HMGCR	$\mu_1,\mu_3,\delta_1,\delta_3$	
Increased Squalene synthase	$\mu_2,\mu_4,\delta_2,\delta_4$	+
Decreased Squalene Synthase	$\mu_2,\mu_4,\delta_2,\delta_4$	
Increased Association of Enzymes	κ_4,κ_5	+
Decreased Association of Enzymes	κ_4,κ_5	
Dissociation of Enzymes	κ_{-4}, κ_{-5}	NC
Increased Product formation	μ_5,μ_7	NC
Decreased Product formation	μ_5	
Decreased Product formation	μ_7	
Increased degrataion of FPP	δ_6	
Decreased degradation of FPP	δ_6	+
Increased degradation	δ_5, δ_7	-
Decreased degradation	δ_5, δ_7	NC
Degradation of cholesterol	δ_8	+ + + /
Stoichiometric coefficients	x_c, x_h, x_s	NC
Genetic binding affinities	K_1, K_2	NC
Increased genetic binding affinity	K_3	NC

Table 3 – continued

Decreased genetic binding affinity	K_3	-
Half Max degradation binding	K_6, K_7, K_8, K_9, K_10	NC
Increased Product formation	μ_6, μ_8	NC
Decreased Product formation	μ_6	-
Decreased Product formation	μ_8	NC
Increased Product formation	μ_9	NC
Decreased Product formation	μ_9	-

483 6. The farnesyl-PP - squalene synthase rate limiting step

Sensitivity analysis of the previous section has revealed evidence of two rate 484 limiting steps working together to regulate homeostatic cholesterol levels. 485 The first is that of the well documented HMGCR HMGCoA reaction, whilst 486 the second involves farmesyl-PP reacting with squalene synthese. Here we 487 investigate the role of the latter reaction, in particular the role of the stoi-488 chometry between farnesyl-PP and squalene synthase in effecting the creation 489 of products downstream of this reaction. In order to do so we consider a sim-490 plified version of this part of the network as given by the reaction stated in 491 equation (47). 492

In this case we have employed A to represent farnesyl-PP, B squalene synthase, C_x the enzyme-substrate complex, P_t squalene and U_a and V_b the influx of substrate and enzyme respectively. For simplicity we assume a constant source of enzyme U_a and substrate V_b , at rates κ_u and κ_v , respectively, and we have removed the effect of the feedback of cholesterol onto squalene synthase degradation. Here κ_{ab} and κ_{-ab} represent the binding and unbinding, respectively, of A and B, κ_p is the rate at which the product is formed and finally

- the degradation of P is represented by δ_{p1} . We observe that $A, B, C_x, P_t \ge 0$ is required for biologically feasible solutions.
- ⁵⁰² Applying the law of mass action to equation (47) leads to

$$\frac{da}{dt} = -2a^2 b\kappa_{ab} + 2c_x \kappa_{-ab} + u_a \kappa_u, \qquad (48)$$

$$\frac{db}{dt} = -a^2 b\kappa_{ab} + c_x \kappa_{-ab} + v_b \kappa_v + c_x \kappa_p, \qquad (49)$$

$$\frac{dc_x}{dt} = a^2 b\kappa_{ab} - c_x \kappa_{-ab} - c_x \kappa_p, \tag{50}$$

$$\frac{dp_t}{dt} = c_x \kappa_p - p_t \delta_{p1}, \tag{51}$$

 $_{\rm 503}$ $\,$ with the initial conditions $\,$

$$a(0) = a_0,$$
 $b(0) = b_0,$ $c_x(0) = 0,$ $p_t(0) = 0.$

We observe that the addition of equations (49) and (50) leads to

$$\frac{da}{dt} + \frac{db}{dt} = v_b \kappa_v,$$

⁵⁰⁴ which for large time becomes

$$b + c_x \sim v_b \kappa_v t. \tag{52}$$

505 This suggests that a, b, c_x and subsequently p_t follow solutions of the form

$$a \sim a_0 t^{\alpha}, \qquad b \sim b_0 t^{\beta}, \qquad c_x \sim c_{x0} t^{\gamma} \quad \text{and} \quad p_t \sim p_{t0} t^{\lambda}.$$
 (53)

Substitution of these solution approximations into equations (48)-(51) leads to

$$a \sim Kt^{-1/2}, \qquad b \sim v_b \kappa_p t, \qquad c_x \sim \frac{u_a \kappa_u}{2\kappa_p} \quad \text{and} \quad p_t \sim \frac{u_a \kappa_u}{2\delta_{p_1}}.$$
 (54)

for which we have the results $a \to 0$, $b \to \infty$ for finite c_x and p_t . This result demonstrates that the substrate farnesyl-PP tends to zero, squalene synthase grows unboundedly in time whilst the complex (bound farnesyl-PP) and product (squalene) remain bounded for any degree of influx.

From this analysis we can conclude that the rate limiting interaction of squa-512 lene synthase and farnesyl-PP would ensure product formation (squalene) is 513 finite and bounded regardless of whether the substrate (farnesyl-PP) or en-514 zyme (squalene synthase) concentrations are bounded. Furthermore, if the 515 levels of squalene are bounded the subsequent products i.e. lanosterol and 516 cholesterol will also be bounded. Thus the mechanism has the downstream 517 effect of ensuring cholesterol levels do not become excessive and alleviates 518 the likelihood of biosynthetic cytotoxicity. 519

⁵²⁰ 7. Steady-state stability analysis

The recent work of Bhattacharya et al. [2] demonstrated that a nonlinear 521 ODE model describing cholesterol biosynthesis via HMGCR mRNA tran-522 scription and subsequent HMGCR translation was monostable. The meval-523 onate pathway examined here is essentially an extension of that model which 524 incorporates further pathway details between HMGCR and cholesterol syn-525 thesis. The increased complexity raises the question as to whether the system 526 exhibits a single real stable steady-state. In this section we utilise a reduction 527 of the full model derived in Section 3 to investigative this. 528

529 7.1. Model reduction

Given the complexity of the governing equations of the full pathway model 530 system (equations (34)-(45)) we begin by simplifying the full pathway (hence-531 forth known as the full model) of Figure 1 by that shown in Figure 3. Here 532 the core product forming and branching points in the pathway have been 533 retained such that w represents SREBP-2, u HMGCR, v squalene synthase, 534 x HMGCoA, y farnesyl-PP and z cholesterol. Here x is produced at a rate 535 A and the negative feedbacks between each relevant component have been 536 included. We further assume that the cholesterol-SREBP-2 negative feed-537 back is the fastest acting process in this reduced network, followed by the 538 synthesis of HMGCR and squalene synthase, which occurs an order of mag-539 nitude slower. Subsequently the formation of x, y and z are assumed to be 540 the slowest in the pathway. Finally, x, y and z decay proportional to their 541 respective concentrations. 542



Figure 3: A model reduction of the mevalonate pathway which incorporates the key enzyme and substrate synthesis processes and branch points, along with their respective feedbacks. Here w represents SREBP-2, u HMGCR, v squalene synthase, x HMGCoA, y farnesyl-PP and z cholesterol, where x is produced at a rate A. It is assumed, as with the full-pathway model, that x, y and z decay proportional to their respective concentrations (not shown here).

⁵⁴³ Applying these assumptions and the law of mass action to the reduced path-⁵⁴⁴ way of Figure 3 leads to the following non-dimensional system of equations

$$\dot{x} = A - \mu_{r1} x u - \delta_{r1} x, \tag{55}$$

$$\dot{y} = \mu_{r1}\beta_x x u - \mu_{r2} y v - \delta_{r2} y, \tag{56}$$

$$\dot{z} = \mu_{r2}\beta_y yv - \delta_{r3} z, \tag{57}$$

$$\epsilon \dot{u} = \mu_{r3}w - \delta_{r4}u \left(\delta_{uz} \frac{z}{\kappa_{r1} + z} + \delta_{uy} \frac{y}{\kappa_{r2} + y} + 1\right), \tag{58}$$

$$\epsilon \dot{v} = \mu_{r4} w - \delta_{r5} v \left(\delta_{vz} \frac{z}{\kappa_{r3} + z} + 1 \right), \tag{59}$$

$$\epsilon^2 \dot{w} = \frac{\alpha_{r1}}{\kappa_{r4} + z^{n_1}} - \delta_{r6} w, \tag{60}$$

 $_{\tt 545}$ $\,$ with the initial conditions $\,$

$$x = 1,$$
 $y = 1,$ $z = 1,$ $u = 1,$ $v = 1$ and $w = 1.$ (61)

Here ϵ represents a small parameter and the remaining model parameters are given by μ_{r1} which represents the rate at which x produces y, μ_{r2} is the rate at which y produces z, μ_{r3} the rate at which u is transcribed, μ_{r4} the rate

at which v is transcribed, β_x and β_y are non-dimensional ratios representing 549 the initial dimensional concentrations of x and y, and y and z, respectively 550 and α_{r1} that rate at which w is produced. The effective binding sites of 551 cholesterol on SREBP-2 is represented by n_1 and $\delta_{r_{1,2,3,4,5,6}}$ represent the 552 rate of degradation of x, y, z, u, v and w, respectively. Finally, $\kappa_{r1,2,3,4}$ are the 553 Michaelis-Menten constants associated with the feedback of z on the rate 554 of u degradation, y on that of u degradation, z on v degradation and z on 555 w inhibition, respectively and δ_{uz} , δ_{uy} and δ_{vz} are dimensionless constants 556 respectively representing their effect. As with the full model of Section 3 557 and for the sake of simplicity we henceforth assume, unless otherwise stated, 558 $\delta_{uz} = 1 = \delta_{uy} = \delta_{vz}.$ 559

Taking the O(1) expansion of equations (55)-(60) leads to

$$\dot{x} = A - \frac{\mu_{r1}x}{(\kappa_{r4} + z^{n_1})\left(\frac{z}{\kappa_{r1} + z} + \frac{y}{\kappa_{r2} + y} + 1\right)} - \delta_{r1}x,$$
(62)
$$\dot{y} = \frac{\tilde{\mu}_{r1}\beta_x x}{(\kappa_{r4} + z^{n_1})\left(\frac{z}{\kappa_{r1} + z} + \frac{y}{\kappa_{r2} + y} + 1\right)} - \frac{\tilde{\mu}_{r2}y}{(\kappa_{r2} + z^{n_1})\left(\frac{z}{\kappa_{r1} + z} + \frac{y}{\kappa_{r2} + y} + 1\right)} - \delta_{r1}x,$$
(62)

$$\dot{y} = \frac{\mu_{r1} \rho_{xx}}{(\kappa_{r4} + z^{n_1}) \left(\frac{z}{\kappa_{r1} + z} + \frac{y}{\kappa_{r2} + y} + 1\right)} - \frac{\mu_{r2} y}{(\kappa_{r4} + z^{n_1}) \left(\frac{z}{\kappa_{r3} + z} + 1\right)} - \frac{\delta_{r2} y}{(\kappa_{r4} + z^{n_1}) \left(\frac{z}{\kappa_{r3} + z} + 1\right)}$$
(63)

$$\dot{z} = \frac{\mu_{r_2} \beta_y y}{\left(\kappa_{r_4} + z^{n_1}\right) \left(\frac{z}{\kappa_{r_3} + z} + 1\right)} - \delta_{r_3} z, \tag{64}$$

561 where

$$\tilde{\mu}_{r1} = \frac{\mu_{r1}\mu_{r3}\alpha_{r1}}{\delta_{r4}\delta_{r6}} \quad \text{and} \quad \tilde{\mu}_{r2} = \frac{\mu_{r2}\mu_{r4}\alpha_{r1}}{\delta_{r5}\delta_{r6}}.$$
(65)

Assuming the concentrations of cholesterol and farnesyl-PP are in excess and the rates of affinity of cholesterol for HMGCR and squalene synthase $(k_{r1} \text{ and } k_{r3})$ and farnesyl-PP for HMGCR are significantly high such that $k_{r1}, k_{r3} \ll z$ and $k_{r2} \ll y$ leads to $k_{r1} + z \sim z$, $k_{r2} + y \sim y$ and $k_{r3} + z \sim z$. Thus

$$\dot{x} = A - \frac{\mu_{rr1}x}{\kappa_{r4} + z^{n_1}} - \delta_{r1}x = f(x, y, z),$$
(66)

$$\dot{y} = \frac{\mu_{rr1}\beta_x x}{\kappa_{r4} + z^{n_1}} - \frac{\mu_{rr2} y}{\kappa_{r4} + z^{n_1}} - \delta_{r2} y = g(x, y, z),$$
(67)

$$\dot{z} = \frac{\mu_{rr2}\beta_y y}{\kappa_{r4} + z^{n_1}} - \delta_{r3} z = h(x, y, z),$$
(68)

567 where $\mu_{rr1} = \frac{1}{3}\tilde{\mu}_{r1}$ and $\mu_{rr2} = \frac{1}{2}\tilde{\mu}_{r2}$.

568 7.2. Steady-state stability

Solving for the steady-states (x^*, y^*, z^*) of equations (66)-(68) leads to the polynomial (recalling that n_1 is an integer)

$$z^{2n_{1}+1}(\delta_{r1}\delta_{r2}\delta_{r3}) + z^{n_{1}+1}(\delta_{r1}\delta_{r3}\mu_{rr2} + 2\delta_{r1}\delta_{r2}\delta_{r3}\kappa_{r4} + \delta_{r2}\delta_{r3}\mu_{rr1}) + z(\delta_{r2}\delta_{r3}\mu_{rr1}\kappa_{r4} + \delta_{r1}\delta_{r2}\delta_{r3}\kappa_{r4}^{2} + \delta_{r1}\delta_{r3}\mu_{rr2}\kappa_{r4} + \delta_{r3}\mu_{rr1}\mu_{rr2}) - A\mu_{rr1}\mu_{rr2}\beta_{x}\beta_{y} = 0, \quad (69)$$

which via Descartes' rule of signs [27] has only one positive root z^* . From (66) and (67) it follows that the corresponding x^* and y^* are also positive.

⁵⁷¹ Now the Jacobian of equations (66)-(68) is given by

$$J = \begin{pmatrix} \frac{-\mu_{rr1}}{\kappa_{r4} + z^{n_1}} - \delta_{r1} & 0 & \frac{n_1 \mu_{rr1} x z^{n_1 - 1}}{(\kappa_{r4} + z^{n_1})^2} \\ \frac{\mu_{rr1} \beta_x}{\kappa_{r4} + z^{n_1}} & \frac{-\mu_{rr2}}{\kappa_{r4} + z^{n_1}} - \delta_{r2} & \frac{-n_1 \mu_{rr1} \beta_x x z^{n_1 - 1}}{(\kappa_{r4} + z^{n_1})^2} + \frac{n_1 \mu_{rr2} y z^{n_1 - 1}}{(\kappa_{r4} + z^{n_1})^2} \\ 0 & \frac{\mu_{rr2} \beta_y}{\kappa_{r4} + z^{n_1}} & \frac{-n_1 \mu_{rr2} \beta_y y z^{n_1 - 1}}{(\kappa_{r4} + z^{n_1})^2} - \delta_{r3} \end{pmatrix},$$

⁵⁷² which allows us to determine the characteristic equation

$$\lambda^3 + a_1\lambda^2 + a_2\lambda + a_3 = 0,$$

573 where

$$a_{1} = -(f_{x} + g_{y} + h_{z}), \qquad a_{2} = f_{x}(g_{y} + h_{z}) + g_{y}h_{z} - g_{z}h_{y}$$

and
$$a_{3} = -(f_{x}(g_{y}h_{z} - g_{z}h_{y}) + f_{z}g_{x}h_{y}).$$
(70)

Now for (x^*, y^*, z^*) to be stable we require $Re(\lambda) < 0$ meaning that the following Routh-Hurwitz conditions [27] must hold

$$a_1 > 0, \quad a_3 > 0 \quad \text{and} \quad a_1 a_2 - a_3 > 0.$$
 (71)

It is easily seen that $f_x, g_y, h_z < 0$ (diagonal entries of J) whilst $f_z, g_x, h_y > 0$. The remaining non-zero term of the Jacobian is

$$g_z = -\left(\frac{n_1 z^{n_1 - 1}}{\kappa_{r4} + z^{n_1}}\right) \left(\frac{\mu_{rr1} \beta_x x}{\kappa_{r4} + z^{n_1}} - \frac{\mu_{rr2} y}{\kappa_{r4} + z^{n_1}}\right)$$

⁵⁷⁸ Both sets of brackets are clearly positive at steady-state (the second set from ⁵⁷⁹ (67)) and so we have $g_z < 0$ at the steady-state.

Using the signs of the Jacobian entries at steady state immediately gives $a_1 > 0$ and, with a little work we can readily use them to deduce that $a_1a_2 - a_3 > 0$. In order to show the remaining required inequality we cannot use the signs of the Jacobian entries alone. Instead we first simplify notation by writing

$$\alpha_1 = \frac{-\mu_{rr1}}{\kappa_{r4} + z^{n_1}}, \qquad \alpha_2 = \frac{-\mu_{rr2}}{\kappa_{r4} + z^{n_1}},$$
$$\gamma_1 = \frac{n_1 \mu_{rr1} x z^{n_1 - 1}}{(\kappa_{r4} + z^{n_1})^2}, \qquad \gamma_2 = \frac{n_1 \mu_{rr2} y z^{n_1 - 1}}{(\kappa_{r4} + z^{n_1})^2},$$

noting that each of these is non-negative. The Jacobian can then be written as

$$J = \begin{pmatrix} -\alpha_1 - \delta_{r1} & 0 & \gamma_1 \\ \alpha_1 \beta_x & -\alpha_2 - \delta_{r2} & -\gamma_1 \beta_x + \gamma_2 \\ 0 & \alpha_2 \beta_y & -\gamma_2 \beta_y - \delta_{r3} \end{pmatrix},$$

587 and a_3 as

$$a_{3} = (\alpha_{1} + \delta_{r1}) ((\alpha_{2} + \delta_{r2})(\gamma_{2}\beta_{y} + \delta_{r3}) + \alpha_{2}\beta_{y}(\gamma_{1}\beta_{x}(\gamma_{1}\beta_{x} - \gamma_{2})) -\gamma_{1}\alpha_{1}\beta_{x}\alpha_{2}\beta_{y} = (\alpha_{1} + \delta_{r1}) (\alpha_{2}\delta_{r3} + \gamma_{2}\beta_{y}\delta_{r2} + \delta_{r2}\delta_{r3}) + \delta_{r1}\alpha_{2}\gamma_{1}\beta_{x}\beta_{y}.$$

Since each symbol is non-negative we immediately have that $a_3 > 0$ as required. Thus (x^*, y^*, z^*) is stable.

In order to provide a check of the stability obtained from the reduced model, we numerically calculated the Jacobian for the full model system using the parameter values detailed in Table 2. All eigenvalues are found to be negative or approximately zero, for a range of initial conditions.

594 8. Periodic solutions

The results of Section 7 have demonstrated that the mevalonate pathway exhibits one real steady-state. Both this model and that of Bhattacharya et ⁵⁹⁷ al. [2] include the negative regulation of SREBP-2 by cholesterol. In the case ⁵⁹⁸ of the three variable model analysed by Bhattacharya and colleagues, they ⁵⁹⁹ demonstrated that the system could exhibit periodic behaviour under certain ⁶⁰⁰ model parameterisations. As such we now investigate numerically whether it ⁶⁰¹ is possible for the mevalonate pathway model to exhibit oscillatory solutions.

⁶⁰² Our investigations focused on the parameters κ_1 , κ_3 , x_c and δ_8 given they ⁶⁰³ are directly involved in the cholesterol-SREBP-2 feedback, are parameters ⁶⁰⁴ for which periodic behaviour was shown in [2], and the results of varying all ⁶⁰⁵ other model parameters in Section 5.1 produced no periodic behaviour.

Local sensitivity analysis of κ_1 , κ_2 , κ_3 , x_c and δ_8 revealed the presence of 606 periodic (damped or undamped) behaviour within the system, an example 607 of which is shown in Figure 4. The presence of oscillatory behaviour for 608 other parameter values showed comparable results. We note the increase in 609 concentration of HMGCoA in Figure 4 is a result of the choice in w made to 610 demonstrate the existence of oscillatory solutions. We sought to numerically 611 investigate further the likelihood of a Hopf bifurication within the mevalonate 612 pathway, as a result of this feedback, and undertook a phase space analysis 613 using MATLAB'S ode15s solver and the plot3 function. We found that the 614 system exhibits an unstable fixed point surrounded by a stable limit cycle 615 and thus appears to undergo a supercritical Hopf bifurcation (results not 616 shown). This was found to be the case when considering the HMGCR mRNA, 617 HMGCR and cholesterol phase space as well as that for squalene synthase 618 mRNA, squalene and cholesterol. 619

These results indicate that the full mevalonate pathway model is able to produce periodic behaviour, similar to that related to more simplified networks within it (e.g. HMGCR mRNA, HMGCR and cholesterol), so long as the global scale negative feedback between cholesterol and SREBP-2 is present.

624 9. Investigating feedbacks

In this section we consider how feedbacks within the mevalonate pathway contribute to the robust control of cholesterol concentrations. Whilst in previous sections we have focused on the role of the global cholesterol-SREBP-2 negative feedback, here we consider the effect of geranyl-PP, farnesyl-PP, lanosterol and cholesterol regulating the degradation of HMGCR, and cholesterol regulating the degradation of squalene synthase, respectively.



Figure 4: Solutions to the system of equations (34) to (46) for which periodic behaviour is exhibited. In this case $\kappa_1 = 1 \times 10^{-12}$, $\kappa_2 = 1 \times 10^{-12}$, $\kappa_3 = 3.74 \times 10^{-4}$, $\delta_8 = 0.1$, with all other parameters as in Table 2.

Given the complexity of the full pathway we began by considering the reduced model shown in Figure 3. This allowed for initial examination of the effect of the feedbacks on the core elements of the network (e.g. rate limiting steps and core products), rather than each individual entity in the full pathway. We identified each feedback in Figure 3 as: (1) $z \rightarrow u$ (cholesterol to HMGCR); (2) $y \rightarrow u$ (farnesyl-PP to HMGCR); and (3) $z \rightarrow v$ (cholesterol to squalene synthase).

We undertook numerical simulations of equations (55) - (60) using the MAT-LAB solver ode15s, assuming $\epsilon = 0.1$ under the eight scenarios detailed in Table 4; when all feedbacks were present, no feedbacks were present, each feedback acted independently and pair-wise. We recorded the difference in steady-state cholesterol concentration, measured as a percentage relative to when all feedbacks were present, in Table 4.

The results in Table 4 clearly show that for fewer feedbacks steady-state 644 cholesterol concentrations increase. When no feedbacks are present, choles-645 terol levels increase by 27.4% in comparison to when all feedbacks are present. 646 Individually, the feedback from farnesyl-PP onto HMGCR has the great-647 est effect on regulating cholesterol levels, whereas those from cholesterol to 648 HMGCR and squalene synthase have the least similar effect. Interestingly 649 our results demonstrate that the feedbacks between cholesterol and HMGCR 650 and squalene synthase, respectively, together have just as tight a control on 651 cholesterol as that of the feedback from farnesyl-PP to HMGCR. The re-652 sults of Table 4 also show that local positive feedbacks affecting the rates of 653 HMGCR and squalene synthase degradation act together with the two rate 654 limiting steps in which they are respectively involved, to tightly regulate the 655 concentration of cholesterol. Importantly, they are able to do so more di-656 rectly and thus more rapidly, given less regulatory steps are involved, than 657 via the feedback between cholesterol and SREBP-2. 658

To test the robustness of the feedback responses, specifically the transient concentration of cholesterol, we introduced a transient influx of cholesterol, B in to z such that

$$B = \begin{cases} 1, & \text{for } 0.10 \le t \le 0.15, \\ 0 & \text{otherwise.} \end{cases}$$
(72)

⁶⁶² under differing feedback scenarios.

Table 4: The percentage relative difference in steady-state cholesterol concentration for when different feedbacks are included compared to when all three feedbacks are in play for the reduced model of Figure 3. In the case of comparing feedbacks either individually or pairwise, the other feedbacks were turned off. Here: (1) $z \to u$ (cholesterol to HMGCR); (2) $y \to u$ (intermediate substrates to HMGCR); and (3) $z \to v$ (cholesterol to squalene synthase) as, defined in Figure 3.

Scenario	Corresponding weighting	Percentage increase in
	parameters	steady-state cholesterol levels.
No feedbacks	$\delta_{uz} = 0 = \delta_{uy} = \delta_{vz}.$	27.4%
(1)	$\delta_{uz} = 1, \delta_{uy} = 0 = \delta_{vz}.$	12.6%
(2)	$\delta_{uz} = 0, \delta_{uy} = 1, \delta_{vz} = 0.$	1.6%
(3)	$\delta_{uz} = 0 = \delta_{uy}, \delta_{vz} = 1.$	12.9%
(1), (2)	$\delta_{uz} = 1 = \delta_{uy}, \delta_{vz} = 0.$	12.6%
(1), (3)	$\delta_{uz} = 1, \delta_{uy} = 0, \delta_{vz} = 1.$	1.6%
(2), (3)	$\delta_{uz} = 0, \delta_{uy} = 1 = \delta_{vz}.$	10.3%



Figure 5: The impact of feedbacks on the reduced model of Figure 3 for the case where z (cholesterol) is increased for 0.10 < t < 0.15. Equations (55)-(60) were solved for all parameter set equal to one with the exception of $\epsilon = 0.1$. Solutions were allowed to reach steady-state before the effect of turning each feedback off was investigated.

Figure 5 demonstrates that each of the feedbacks tightly regulate the concen-663 trations of x, y and z and that varying combinations of them did not alter the 664 overall transient behaviour. Additionally in scenarios where feedback (3) was 665 turned off, levels of y (farnesyl-PP) were very low as more squalene synthase 666 is available to bind with farnesyl-PP to form squalene. This coupled with 667 the analysis undertaken in Section 6 showing if the concentration of squalene 668 synthase grows unbounded the rate limiting step between it and farnesyl-PP 669 acts to control the downstream concentrations of lanosterol and subsequently 670 cholesterol, demonstrates these two processes act together locally to tightly 671 regulate cholesterol levels in this section of the pathway. 672

We undertook the same analysis of each feedback on the full model of the 673 pathway, equations (34) to (46). We inhibited the feedbacks from: (1) choles-674 terol to HMGCR degradation; (2) farnesyl-PP to HMGCR degradation; and 675 (3) cholesterol to squalene synthase degradation. We again conducted the 676 same eight scenarios detailed in Table 4 and found all scenarios show the same 677 transient behaviour in good agreement with the reduced model. The only 678 notable change was were switching feedback (2) off led to slightly higher lev-679 els of HMGCR. This difference was not seen when feedback (3) was switched 680 on concurrently to feedback (2). 681

⁶⁸² 10. CYP51 inhibition

So far we have demonstrated that cholesterol biosynthesis via the mevalonate 683 pathway is a tightly regulated process; a result of two enzymatic rate limiting 684 steps coupled with local and global feedbacks within the signalling network. 685 In this section we show how these elements integrate together to ensure a 686 robust network response to the effect of the fungicide agent CYP51. CYP51 687 is known to inhibit post lanosterol production processes and is used in crop 688 protection as an anti-fungal agent. It acts by reducing cholesterol concen-689 trations within the cell, thereby compromising cell wall integrity, ultimately 690 leading to cell death. Concerns exist that this inhibition is likely to lead to 691 increases in farnesyl-PP levels, thereby inducing unwanted side-effects within 692 other cell signalling cascades who share cross-talk with farnesyl-PP. 693

To investigate the effect of CYP51 inhibition on the pathway we first ran the system of equations (34) to (46) to steady-state. Taking this as our starting



Figure 6: The effect of CYP51 inhibition on equations (34) to (46). Here $\mu_9 = 0$ at $\bar{t} = 5$ hours for 2 hours with $\kappa_3=0.075$, to simulate CYP51 inhibition as described by equation (73). Cholesterol concentrations decline which leads to a decrease in HMGCR levels as a result of the feedback between cholesterol and SREBP-2. Hence concentrations of geranyl-PP, farnesyl-PP and squalene all decline. All concentrations return to steady-state after CYP51 inhibition stops.

⁶⁹⁶ point we then simulated the effect of CYP51 inhibitors by letting

$$\mu_9 = \begin{cases} 0, & \text{for } 5 \le t \le 7, \\ 3.61 \times 10^2 & \text{otherwise.} \end{cases}$$
(73)

Results are shown in Figure 6. We see that CYP51 inhibition leads to a 697 sharp increase in lanosterol and decline in cholesterol concentrations. Here 698 we would expect an increase in HMGCR concentrations due to the rise in 699 HMGCR mRNA, however the sharp increase in lanosterol concentration 700 causes the degradation of HMGCR to be up-regulated, and so its concen-701 tration subsequently declines. The reduction in HMGCR thus leads to a 702 decline in the central cascade products of geranyl-PP, farnesyl-PP and squa-703 lene. As a result we see that the change in these central cascade products is 704 limited and that faresyl-PP levels actually reduce when production of choles-705 terol from lanosterol is inhibited. We note that an increase in inhibition of 706 SREBP-2 by cholesterol ($\kappa_3 = 0.075$) was required in order to observe a 707 response in HMGCR mRNA and squalene synthase mRNA. 708

⁷⁰⁹ 11. Summary and conclusions

We have formulated, parameterised and analysed a nonlinear ODE model of the mevalonate cholesterol biosynthesis pathway. Our results show that the pathway tightly regulates steady-state and transient cholesterol levels via two rate limiting steps, internal local positive feedbacks affecting the rate of degradation of certain products within the pathway and a global negative feedback between cholesterol and SREBP-2.

A local sensitivity analysis of the model revealed a number of important reg-716 ulatory points within the pathway. It highlighted that decreases in HMGCR 717 levels has the greatest impact on downstream cholesterol levels either via 718 variation in transcription or translation rates or the rate of HMGCR mRNA 719 or HMGCR degradation. Increasing products prior to farnesyl-PP interact-720 ing with squalene synthase has a more significant effect on cholesterol levels 721 in contrast to those after the reaction, the rates at which geranyl-PP and 722 squalene are formed have the most significant effect. Altering the rate of 723 cholesterol esterification has a significant impact on HMGCR and squalene 724 synthase levels via the cholesterol SREBP-2 negative feedback loop. 725

Our sensitivity analysis also revealed the importance of the rate limiting en-726 zyme substrate reactions of HMGCoA with HMGCR and farnesyl-PP with 727 squalene synthase, the latter augmented by separate analytical analysis of the 728 farnesyl-PP squalene synthase rate limiting step. The HMGCR-HMGCoA 729 reaction was found to be an important upstream regulator of all main path-730 way products. That of farnesyl-PP and squalene synthase was found to 731 be important in not only regulating downstream production of squalene, 732 lanosterol and thus cholesterol, but in ensuring their levels did not increase 733 significantly if levels of farnesyl-PP and squalene synthase did. 734

Analysis of a reduced model of the full pathway, which captured the main 735 products and interactions between them, demonstrated that the system ex-736 hibits one real stable steady-state. The global feedback between cholesterol 737 and SREBP-2 leads to monotonic, oscillatory and damped oscillatory be-738 haviour, which agrees with the simplified HMGCR cholesterol regulatory 739 model of [2]. This result shows that the feedback between cholesterol and 740 SREBP-2 acts to globally regulate the dynamic pathway behaviour. This 741 is in contrast to internal positive feedbacks between geranyl-PP, farnesylPP, 742 lanosterol and the degradation of HMGCR and squalene synthese which our 743 analysis demonstrated act directly within the pathway to tightly regulate 744 overall cholesterol concentrations. 745

It is clear that feedbacks in the pathway act to control the dynamical re-746 sponse, enzyme concentrations and hence the concentration of cholesterol. 747 The cholesterol-SREBP-2 feedback allows for cholesterol regulation of its own 748 production over a longer timescale than those from geranyl-PP, farnesyl-PP, 749 lanosterol and cholesterol to HMGCR and cholesterol to squalene synthase; 750 which respond directly within the pathway to any variation in cholesterol lev-751 els. These direct responses alleviate the effect of further reactions in delaying 752 the reduction of the entity they are targeting in the pathway. 753

Further evidence of the system's robust network control via the integration 754 of two rate limiting steps and feedbacks was shown in the case of CYP51 755 inhibition. Simulations of CYP51 inhibition show the network response pre-756 vents cytotoxic build up of central cascade products geranyl-PP, squalene and 757 farnesyl-PP. This is important since increased farnesyl-PP levels are linked 758 with several other signalling pathways and excessive amounts are thought 759 to cause tumours. In this way we have shown that CYP51 inhibitors would 760 have little effect on farnesyl-PP concentrations in the mevalonate pathway. 761

Given the importance of cholesterol synthesis in maintaining the integrity of 762 cell function for many cellular phenotypes, the results of the work here are in 763 many ways unsurprising. Cholesterol levels need to be tightly regulated, both 764 in response to internal cellular variations and external factors, e.g. disease or 765 dietary factors. Our work here has clearly demonstrated that the pathway is 766 robustly designed and includes a number of 'fail safe' type mechanisms in the 767 form of regulatory feedbacks and rate limiting steps which act in concert to 768 provide a robust regulatory system. These results are in agreement with the 769 work of August et al. [1] and Morgan et al. [25], who both demonstrated that 770 the cholesterol biosynthesis aspects of their models were robust to parameter 771 variation. The design of the network ensures that the integrity of cholesterol 772 levels is not greatly compromised, should one or more of these mechanisms 773 fail, thus ensuring cell survival is maintained. 774

775 Acknowledgement

FP acknowledges the support of a Engineering Physical Sciences Research
Council (EPSRC) UK CASE studentship in collaboration with Syngenta
(EP/P505682/1 & EP/J500501/1). MJT is grateful for the support of a
Research Council UK Fellowship (EP/C508777/1) during parts of the period
in which this work was undertaken.

⁷⁸¹ Appendix A. Biochemical reaction details

In order to formulate a mathematical model of the interactions shown in
Figure 1 we first consider the biochemical details of each reaction. The
binding of SREBP-2 to HMGCR DNA and subsequent mRNA and protein
formation is governed by

$$G_h + x_h S \xrightarrow{\bar{\kappa}_1} S_{bh} \xrightarrow{\bar{\mu}_1} M_h \xrightarrow{\bar{\mu}_3} H_r,$$
 (A.1)

where HMGCR free DNA is represented by G_h , S is SREBP-2, S_{bh} is SREBP-2 bound to the DNA, M_h is HMGCR mRNA and H_r is HMGCR. The constant reaction rates $\bar{\kappa}_1$ and $\bar{\kappa}_{-1}$ represent the binding and unbinding of SREBP-2 and DNA protein respectively, $\bar{\mu}_1$ is the rate of transcription of HMGCR mRNA and $\bar{\mu}_3$ is the rate of HMGCR translation. Finally x_h is the number of binding sites on the DNA that SREBP-2 must bind to.

⁷⁹² Binding of SREBP-2 to squalene synthase DNA and subsequent mRNA and
 ⁷⁹³ protein formation is governed by

$$G_{ss} + x_s S \xrightarrow{\bar{\kappa}_2} S_{bss} \xrightarrow{\bar{\mu}_2} M_{ss} \xrightarrow{\bar{\mu}_4} S_s,$$
 (A.2)

where free DNA binding sites responsible for squalene synthesis is 794 represented by G_{ss} , S_{bss} is SREBP-2 bound to the DNA, M_{ss} is squalene 795 synthase mRNA and S_s is squalene synthase. The constant reaction rates 796 $\bar{\kappa}_2$ and $\bar{\kappa}_{-2}$ represent the binding and unbinding of SREBP-2 and DNA re-797 spectively, $\bar{\mu}_2$ is the rate of transcription of mRNA responsible for squalene 798 synthase and $\bar{\mu}_4$ is the rate of translation of squalene synthase from mRNA. 799 Finally x_s is the number of binding sites on the DNA that SREBP-2 must 800 bind to. 801

Binding of HMGCR and HMGCoA and subsequent production of geranyl-PP
 and farnesyl-PP is governed by

where free HMGCoA is represented by H_c , H_b is HMGCR bound to HMG-CoA, G_{pp} is geranyl-PP and F_{pp} is farnesyl-PP. The constant reaction rates $\bar{\kappa}_4$ and $\bar{\kappa}_{-4}$ represent binding and unbinding of HMGCR and HMGCoA respectively, $\bar{\mu}_5$ is the rate of production of geranyl-PP and $\bar{\mu}_6$ is the rate of production of farnesyl-PP.

Two molecules of farnesyl-PP bind to one molecule of squalene synthase for the subsequent production of squalene, lanosterol and cholesterol such that

$$2F_{pp} + S_s \xrightarrow{\bar{\kappa}_5} F_{bpp} \xrightarrow{\bar{\mu}_7} S_q + S_s$$
$$\bar{\mu}_8 \downarrow \qquad (A.4)$$
$$L \xrightarrow{\bar{\mu}_9} C,$$

where bound farnesyl-PP and squalene synthase is represented by F_{bpp} , S_q is squalene, L is lanosterol and C is cholesterol. The constant reaction rates $\bar{\kappa}_5$ and $\bar{\kappa}_{-5}$ denote binding and unbinding of squalene synthase and farnesyl-PP respectively, $\bar{\mu}_7$ is the rate of squalene production, $\bar{\mu}_8$ is the rate of lanosterol production $\bar{\mu}_9$ that of cholesterol.

⁸¹⁶ The negative regulation of SREBP-2 by cholesterol is governed by

$$x_c C + S \xrightarrow[\bar{\kappa}_{3}]{\bar{\kappa}_{-3}} C_b,$$
 (A.5)

where bound cholesterol and SREBP-2 is represented by C_b , the constant reaction rates $\bar{\kappa}_3$ and $\bar{\kappa}_{-3}$ represent the binding and unbinding of cholesterol and SREBP-2, respectively. Finally x_c is the number of binding sites that must be occupied by cholesterol on SREBP-2 to inactivate SREBP-2.

⁸²¹ Each degradation process is described by

where δ_i $(i \in [1, ..., 8])$ are the rates of degradation of each mRNA, protein and enzyme, respectively.

Appendix B. Model reduction

We begin by observing three conservation relations. Firstly, the total amount of DNA within a cell remains constant such that

$$\bar{g}_h + \bar{s}_{bh} = \bar{g}_{h0}$$
 and $\bar{g}_{ss} + \bar{s}_{bss} = \bar{g}_{ss0}$, (B.1)

which are formed from the addition and integration (with respect to time) of equations (1) and (4), and (2) and (5), respectively.

⁸²⁹ The total amount of SREBP-2 in a cell is also constant which similarly gives

$$\bar{s} + \bar{c}_b = \bar{s}_0,\tag{B.2}$$

using equations (3) and (18).

We assume the following reactions occur on a faster timescale than others in the signalling cascade and as such invoke the quasi-steady-state approximation. We assume DNA-transcription factor binding is rapid [20, 2] such that from equation (4)

$$\bar{\kappa}_1 \bar{s}^{x_h} (\bar{g}_{h0} - \bar{s}_{bh}) - \bar{\kappa}_{-1} \bar{s}_{bh} \approx 0,$$

where we have substituted for \bar{g}_h using the first conservation relationship in equation (B.1). This result can be re-arranged for $s_{\bar{b}h}$ to give

$$\bar{s}_{bh} \approx \frac{\bar{g}_{h0}\bar{s}^{x_h}}{\bar{s}^{x_h} + \bar{K}_1^{x_h}},\tag{B.3}$$

with $\bar{K}_1 = \left(\frac{\bar{\kappa}_{-1}}{\bar{\kappa}_1}\right)^{\frac{1}{x_h}}$.

Using the second conservation relationship in equation (B.1) and applying the same assumption to equation (5) yields

$$\bar{s}_{bss} \approx \frac{\bar{g}_{ss0}\bar{s}^{x_s}}{\bar{s}^{x_s} + \bar{K}_2^{x_s}},\tag{B.4}$$

with $\bar{K}_2 = \left(\frac{\bar{\kappa}_{-2}}{\bar{\kappa}_2}\right)^{\frac{1}{x_s}}$.

Finally we assume that cholesterol-SREBP-2 binding is also rapid such that from equation (3)

$$\bar{s} \approx \frac{\bar{K}_3^{x_c} \bar{s}_0}{\bar{c}^{x_c} + \bar{K}_3^{x_c}} = \frac{\bar{s}_0}{1 + (\frac{\bar{c}}{\bar{K}_3})^{x_c}},\tag{B.5}$$

with $\bar{K}_3 = \left(\frac{\bar{\kappa}_{-3}}{\bar{\kappa}_3}\right)^{\frac{1}{x_c}}$. This relationship can subsequently be used to express \bar{s}_{bh} and \bar{s}_{bss} in terms of c.

Using the results of equations (B.3), (B.4) and (B.5) we can simplify equations (6) and (7) to

$$\frac{d\bar{m}_h}{d\bar{t}} = \frac{\bar{\mu}_1^*}{1 + \left(\frac{\bar{K}_1(1 + (\frac{\bar{c}}{\bar{K}_3})^{x_c})}{\bar{s}_0}\right)^{x_h}} - \bar{\delta}_1 \bar{m}_h, \tag{B.6}$$

843 and

$$\frac{d\bar{m}_{ss}}{d\bar{t}} = \frac{\bar{\mu}_2^*}{1 + \left(\frac{\bar{K}_2(1 + (\frac{\bar{c}}{\bar{K}_3})^{x_c})}{\bar{s}_0}\right)^{x_s}} - \bar{\delta}_2 \bar{m}_{ss},\tag{B.7}$$

844 where $\bar{\mu}_1^* = \bar{\mu}_1 \bar{g}_{h0}$ and $\bar{\mu}_2^* = \bar{\mu}_2 \bar{g}_{ss0}$.

Equation (31) is derived from equations (3), (4), (5) and (17), respectively, such that $\bar{z} = \bar{z}$

$$\frac{d}{d\bar{t}}(\bar{s} + x_h\bar{s}_{bh} + x_s\bar{s}_{bss} - \bar{c}/x_c) = \frac{\bar{\mu}_9\bar{t} - \bar{\delta}_8\bar{c}}{x_c}$$

which leads to

$$(1 - x_c(\bar{s}' + x_h\bar{s}'_{bh} + x_s\bar{s}'_{bss}))\frac{d\bar{c}}{d\bar{t}} = \bar{\mu}_9\bar{l} - \bar{\delta}_8\bar{c},$$

via the chain rule, where ' denotes differentiation with respect to \bar{c} such that from (B.5), (B.3) and (B.4) we have

$$\frac{d\bar{s}}{d\bar{c}} = \frac{-\bar{s}_0 x_c \left(\frac{\bar{c}}{\bar{K}_3}\right)^{x_c}}{\bar{c} \left(1 + \left(\frac{\bar{c}}{\bar{K}_3}\right)^{x_c}\right)^2},\tag{B.8}$$

847

$$\frac{d\bar{s}_{bh}}{d\bar{c}} = \frac{d\bar{s}_{bh}}{d\bar{s}}\frac{d\bar{s}}{d\bar{c}} = \frac{x_h\bar{g}_{h0}\bar{K}_1^{x_h}\bar{s}^{x_h-1}}{(\bar{s}^{x_h}+\bar{K}_1^{x_h})^2}\frac{d\bar{s}}{d\bar{c}}$$
(B.9)

848 and

$$\frac{d\bar{s}_{bss}}{d\bar{c}} = \frac{d\bar{s}_{bss}}{d\bar{s}}\frac{d\bar{s}}{d\bar{c}} = \frac{x_s\bar{g}_{ss0}\bar{K}_2^{x_s}\bar{s}^{x_s-1}}{(\bar{s}^{x_s} + \bar{K}_2^{x_s})^2}\frac{d\bar{s}}{d\bar{c}},\tag{B.10}$$

respectively. Here \bar{g}_{h0} and \bar{g}_{ss0} are the total concentration of HMGCR and squalene synthase DNA, respectively, in a cell.

⁸⁵¹ Appendix C. Parameter details

In this section we detail, where relevant, calculations used to estimate the parameters detailed in Table 1.

⁸⁵⁴ $\bar{\mathbf{m}}_{\mathbf{h0}}$ - Initial concentration of HMGCR mRNA: Ruddling et al. [30] ⁸⁵⁵ details copy numbers of mRNA found in human liver cells under basal con-⁸⁵⁶ ditions. So we take a value of 30 copies of HMGCR mRNA per cell i.e. per ⁸⁵⁷ 10^{-9} ml. So

 $\frac{30 \text{ molecules}}{1 \times 10^{-9} \text{ml}} = 3.0 \times 10^{10} \text{ molecules/ml}.$

This value was then refined using local sensitivity analysis to give $\bar{m}_{h0} = 3.0 \times 10^9$ molecules/ml.

⁸⁶⁰ $\bar{\mathbf{m}}_{ss0}$ - Initial concentration of squalene synthase mRNA: Ruddling ⁸⁶¹ et al. [30] details copy numbers of mRNA found in human liver cells under ⁸⁶² basal conditions. So we take a value of 30 copies of squalene synthase mRNA ⁸⁶³ per cell i.e. per 10⁻⁹ ml. So

 $\frac{30 \text{ molecules}}{1 \times 10^{-9} \text{ml}} = 3.0 \times 10^{10} \text{molecules/ml}.$

This value was then refined using local sensitivity analysis to give $\bar{m}_{ss0} = 3.0 \times 10^9$ molecules/ml.

 $\bar{\mathbf{s}}_{sT}$ - Total concentration of squalene synthase: One liver cell contains 300pg/cell protein and has a volume of 10^{-9} ml. Bruenger and Rilling [5] state there are 4.2 nmol of squalene synthase per gram of wet tissue such that

 4.2×10^{-9} mol/g tissue $\times 6.022 \times 10^{23}$ molecules/mol

870 which gives

$$\frac{2.53 \times 10^{15} \text{molecules } /\text{g} \times 1.00 \times 10^{-12}}{10^{-9} \text{ml}} = 7.59 \times 10^{14} \text{ molecules /ml.}$$

 $\bar{\mathbf{h}}_{cT}$ - Total concentration of HMGCoA: One liver cell contains approximately 300pg/cell protein and has volume 10^{-9} ml/cell. The molecular weight of HMGCoA is 199.659 g/mol according to human metabolic database [38]. Then we know

$$\frac{300 \times 10^{-12} \text{g}}{199.659 \text{g/mol}} = 3.29 \times 10^{-13} \text{mol/cell}.$$

So we have, per cell, $3.29 \times 10^{-13} \text{mol}/10^{-9} \text{ml} = 3.92 \times 10^{-4} \text{mol/ml}$. Applying Avagadro's number we can find the number of molecules per ml

 $3.92 \times 10^{-4} \text{ mol/ml} \times 6.022 \times 10^{23} \text{ molecules/mol} = 1.98 \times 10^{20} \text{ molecules/ml}.$

Segel (1993) [33] states a cell contains an average of 1000 enzymes, so we have 9.04 × 10¹⁴ molecules/ml. This value was then refined using local sensitivity analysis to give $\bar{h}_{cT} = 1.98 \times 10^{15}$ molecules/ml.

 $\bar{\mathbf{g}}_{\mathbf{h0}}, \bar{\mathbf{g}}_{\mathbf{ss0}}$ - **HMGCR and squalene synthase gene concentration**: The molecular weight of the HMGCR gene is 97,476 Da [41], whilst that of the human genome is 2 ×10¹²Da [42]. The total quantity of DNA in a cell weighs 7pg, such that that of HMGCR is 3.41×10^{-7} pg. Observing that 1 Da is equivalent to 1g/mol and assuming the volume of a cell is 1 nml, we have

$$\frac{3.41 \times 10^{-7} \text{ pg} \times 6.023 \times 10^{23} \text{ molecules/mol}}{97,476 \text{ g/mol} \times 1 \text{ nml}} = 2.11 \times 10^9 \text{molec/ml}.$$

We likewise assume the squalene synthase gene (with no further details available) is the same concentration.

⁸⁸⁷ $\bar{\mu}_1^*$ - Rate of HMGCR mRNA transcription: Darzacq et al. [8] states ⁸⁸⁸ 12 bases are transcribed per second. Goldstein and Brown [12] say one ⁸⁸⁹ HMGCoA-R gene is 24826 bases long. Therefore we have

$$\frac{24826 \text{ bases}}{12 \text{ bases/s}} = 2068.83 \text{s}.$$

We add 30 minutes to account for post transcriptional processing steps of mRNA cleavage giving 3868.83s. So for one gene we have

$$\frac{1 \text{ molecule}}{3868.83 \text{s}} = 2.58 \times 10^{-4} \text{ molecules/s}.$$

A liver cell is somatic and hence diploid meaning it contains contains two genes, so we have

$$2.58 \times 10^{-4}$$
 molecules/s $\times 2 = 5.17 \times 10^{-4}$ molecules/s.

The average cell volume is $1pl = 1 \times 10^{-9}ml$ so the rate of transcription is given by

$$\frac{5.17 \times 10^{-4} \text{ molecules/s}}{1 \times 10^{-9} \text{ml}}$$

giving $\bar{\mu}_1^* = 5.17 \times 10^5$ molecules/ml/s.

⁸⁹⁷ $\bar{\mu}_2^*$ - Rate of squalene synthase mRNA transcription: Darzacq et al. [8] ⁸⁹⁸ states 12 base pairs are transcribed per second. Tansey & Shechter [37] say ⁸⁹⁹ one human squalene synthase gene is over 30000 bases long. Therefore we ⁹⁰⁰ have

$$\frac{30000 \text{ bases}}{12 \text{ bases/s}} = 2500 \text{s}.$$

We add 30 minutes to account for post transcriptional processing steps of mRNA cleavage giving 4300 thus for one gene we have

$$\frac{1 \text{ molecule}}{4300 \text{s}} = 2.33 \times 10^{-4} \text{ molecules/s}.$$

A liver cell is somatic and hence diploid meaning it contains contains two
 genes, so we have

$$2.33 \times 10^{-4}$$
 molecules/s $\times 2 = 4.65 \times 10^{-4}$ molecules/s.

The average cell volume is $1pl = 1 \times 10^{-9}ml$ so the rate of transcription is given by

$$\frac{4.65 \times 10^{-4} \text{ molecules/s}}{1 \times 10^{-9} \text{ml}}$$

giving $\bar{\mu}_2^* = 4.65 \times 10^5$ molecules/ml/s.

⁹⁰⁸ $\bar{\mu}_3$ - Rate of HMGCR translation: Trachsel [39] states 6 amino acids are ⁹⁰⁹ translated per second. One amino acid is encoded by 3 bases or nucleotides. ⁹¹⁰ HMGCR mRNA transcript has 4475 bases (Goldstein & Brown [12]). Hence ⁹¹¹ transcription takes:

$$\frac{4475 \text{ bases}}{6 \text{ amino acids/s} \times 3 \text{ amino acids/base}} = 248.61s,$$

⁹¹² We add 60 minutes to account for the initiation of this process

⁹¹³ Then per ribosome we have

$$\frac{1 \text{ molecule}}{3848.61 \text{s}} = 2.60 \times 10^{-4} \text{ molecules/s/ribosome.}$$

- ⁹¹⁴ A ribosome can only attach every 35 bases do to its size meaning 1 mRNA
- ⁹¹⁵ molecule has 127.86 ribosomes attached.
- ⁹¹⁶ Then per mRNA molecule we have:

 2.60×10^{-4} molecules/s/ribosome \times 127.86 ribosomes/molecule

917 giving $\bar{\mu}_3 = 3.32 \times 10^{-2}$ /s.

⁹¹⁸ $\bar{\mu}_4$ - Rate of squalene synthase translation: Trachsel [39] states 6 amino ⁹¹⁹ acids are translated per second. One amino acid is encoded by 3 bases ⁹²⁰ or nucleotides. Jiang et al. [36] state that one squalene synthase mRNA ⁹²¹ transcript contains 2502 bases. Hence transcription takes:

 $\frac{2502 \text{ bases}}{6 \text{ amino acids/s} \times 3 \text{ amino acids/base}} = 139 \text{s},$

⁹²² We add 60 minutes to account for the initiation of this process

3739s.

⁹²³ Then per ribosome we have

$$\frac{1 \text{ molecule}}{3739 \text{s}} = 2.67 \times 10^{-4} \text{ molecules/s/ribosome.}$$

- A ribosome can only attach every 35 bases do to its size meaning 1 mRNA molecule has 71.49 ribosomes attached.
- ⁹²⁶ Then per mRNA molecule we have:

 2.67×10^{-4} molecules/s/ribosome \times 71.49 ribosome/molecule

927 giving $\bar{\mu}_4 = 1.91 \times 10^{-2}$ /s.

⁹²⁸ $\bar{\mu}_5$ - Rate of geranyl-PP synthesis: Tanaka et al. [47] tell us that liver ⁹²⁹ microsomes form 52 pmol mevalonate per minute per mg protein. Istvan et ⁹³⁰ al. [15] say HMGCR is tetrameric arranged in 2 dimer, with 4 active sites, ⁹³¹ has molecular weight 199812 Da. The activity of the enzyme is where

$$52 \times 10^{-12} \text{mol/min/mg protein} \approx 52 \times 10^{-12} \times N_A.$$

 $_{932}$ $N_A = 6.022 \times 10^{23}$ is Avagadro's constant. So we have

 52×10^{-12} mol/min/mg protein $\times 6.022 \times 10^{23}$ molecules/mol

 $= 3.13 \times 10^{13}$ molecules/min/mg protein.

Segel [33] says there's 1000 different enzymes in a cell, so for 1 mg of protein we have

$$\frac{1 \times 10^{-3} \text{g}}{199812 \text{g/mol} \times 1000} = 5.00 \times 10^{-12} \text{mol}.$$

Given there are 4 active sites per HMGA-CoA Reductase enzyme, there are 2.00×10⁻¹¹ moles of enzyme active sites in 1 mg of protein. Given the specific activity of an enzyme we find $\bar{\mu}_5$ is equal to

$$\frac{52 \times 10^{-12} \text{ mol/min/mg}}{2.00 \times 10^{11} \text{mol/mg}} = 2.60 \text{min}$$

939 giving $\bar{\mu}_5 = 4.33 \times 10^{-2}$ /s.

⁹⁴⁰ $\bar{\mu}_6$, $\bar{\mu}_8$ and $\bar{\mu}_9$ - Rates of farnesyl-PP, lanosterol and cholesterol syn-⁹⁴¹ thesis: Since the value for $\bar{\mu}_5$ is used to describe cholesterol production from ⁹⁴² HMGCR, we can assume all steps in between must occur at the same rate ⁹⁴³ or faster. Therefore we set $\bar{\mu}_6$, $\bar{\mu}_8$ and $\bar{\mu}_9$ equal to 4.33×10^{-2} /s.

⁹⁴⁴ $\bar{\mu}_7$ - Rate of squalene synthesis. Since the value for $\bar{\mu}_5$ is used to describe ⁹⁴⁵ cholesterol production from HMGCR, we can assume all steps in between ⁹⁴⁶ must occur at the same rate or faster. Therefore as an estimate we set $\bar{\mu}_7$ ⁹⁴⁷ equal to 4.33×10^{-2} /s. This value was then refined using local sensitivity ⁹⁴⁸ analysis to give $\bar{\mu}_7 = 2.17 \times 10^{-1}$ 1/s.

⁹⁴⁹ $\overline{\mathbf{K}}_{1}$ - Disassociation constant of SREBP-2 for HMGCR DNA: Yang ⁹⁵⁰ and Swartz [29] quantifed DNA binding affinities to other transcription fac-⁹⁵¹ tors at 54.2 nmol. We convert this value into units of molecules/ml by the ⁹⁵² use of Avogadro's constant.

$$\frac{100 \times 10^{-9} \text{ moles}}{1000 \text{ml}} \times 6.022 \times 10^{23} \text{ molecules/mol} = 3.26 \times 10^{13} \text{ molecules/ml}.$$

This value was then refined using local sensitivity analysis to give $\bar{K}_1 = 8.21 \times 10^{12}$ molecules/ml.

⁹⁵⁵ $\overline{\mathbf{K}}_2$ - Disassociation constant of SREBP-2 for squalene synthase ⁹⁵⁶ DNA: This was assumed equivalent to that of SREBP-2 for HMGCR DNA, ⁹⁵⁷ i.e. 3.26×10^{13} molecules/ml. The value was then refined using local sensi-⁹⁵⁸ tivity analysis to give $\overline{K}_2 = 8.21 \times 10^{12}$ molecules/ml.

933

 $\bar{\mathbf{K}}_{3}$ - Disassociation constant of SREBP-2 for cholesterol: Radhakrishnan et al. [46] state the binding reaction between cholesterol and SCAP is saturable and half-maximal binding occurs at approximately 100 nmol. We convert this value into units of molecules/ml by the use of Avogadro's constant.

 $\frac{100 \times 10^{-9} \text{ moles}}{1000 \text{ ml}} \times 6.022 \times 10^{23} \text{ molecules/mol} = 6.02 \times 10^{13} \text{ molecules/ml},$

as an estimate we took $\bar{K}_3 = O(10^{14})$. This value was then refined using local sensitivity analysis to give $\bar{K}_3 = 1.49 \times 10^{16}$ molecules/ml.

⁹⁶⁶ $\bar{\kappa}_4$ and $\bar{\kappa}_{-4}$ - Forward and reverse rates of HMGCR binding to HMG-⁹⁶⁷ CoA: These values were initially informed by assuming the ratio of $\bar{\kappa}_4/\bar{\kappa}_{-4}$ ⁹⁶⁸ were the same order as those of \bar{K}_1 , \bar{K}_2 and \bar{K}_3 . We then assumed $\bar{\kappa}_{-4} \ll \bar{\kappa}_4$ ⁹⁶⁹ whereby we took an initial estimate of $\bar{\kappa}_{-4} = 1 \times 10^{-3}$ /s. These values were ⁹⁷⁰ then adjusted, via a sensitivity analysis, to give the required steady-state ⁹⁷¹ cholesterol levels. This resulted in values of $\bar{\kappa}_4 = 1.39 \times 10^{-16}$ ml/molecules ⁹⁷² s and $\bar{\kappa}_{-4} = 1.75 \times 10^{-7}$ /s.

⁹⁷³ $\bar{\kappa}_5/\bar{\kappa}_5$ - Forward and reverse rates of farnesyl-PP binding to squalene ⁹⁷⁴ synthase: These values were obtained in a similar manner to those of $\bar{\kappa}_4$ ⁹⁷⁵ and $\bar{\kappa}_{-4}$. This led to $\bar{\kappa}_5 = 1.76 \times 10^{-30}$ ml/molecule s and $\bar{\kappa}_{-5} = 1.75 \times 10^{-5}$ ⁹⁷⁶ /s.

⁹⁷⁷ $\bar{\mathbf{K}}_{6}$, $\bar{\mathbf{K}}_{7}$, $\bar{\mathbf{K}}_{8}$, $\bar{\mathbf{K}}_{9}$ and $\bar{\mathbf{K}}_{10}$ - Michaelis-Menten constants of geranyl-PP, ⁹⁷⁸ farnesyl-PP, lanosterol and cholesterol for HMGCR degradation ⁹⁷⁹ and cholesterol for squalene synthase degradation, respectively: ⁹⁸⁰ These were determined as the half-maximal values which produced a sig-⁹⁸¹ moidal type response for each of the respective cascade products.

⁹⁸² $\bar{\delta}_1$ - Degradation rate of HMGCR mRNA. Degradation rates of proteins ⁹⁸³ and mRNAs are based on their half lives, derived from an exponential decay ⁹⁸⁴ model. Wilson and Deeley [3] state HMGCR mRNA has a half life of 4.3 ⁹⁸⁵ hours, measured in Hep G2 cells, giving $\bar{\delta}_1$ =ln 2/15480s=4.48×10⁻⁵/s.

 $\bar{\delta}_2$ - Degradation rate of squalene synthase mRNA: This was assumed equivalent to that of HMGCR mRNA.

⁹⁸⁸ $\bar{\delta}_3$ - Degradation of HMGCR: Brown et al. [44] found HMGCR protein ⁹⁸⁹ has a half life of 3 hours, measured in human fibroblast cells, such that ⁹⁹⁰ $\bar{\delta}_3 = \ln 2/10800s = 6.42 \times 10^{-5}/s.$

- ⁹⁹¹ $\overline{\delta}_4$ Squalene synthase degradation rate: This was assumed equivalent ⁹⁹² to that of HMGCR.
- ⁹⁹³ $\bar{\delta}_5$, $\bar{\delta}_6$ and $\bar{\delta}_7$ Degradation rates of geranyl-PP, farnesyl-PP and ⁹⁹⁴ lanosterol: These were assumed equivalent to that of cholesterol.
- ⁹⁹⁵ $\overline{\delta}_8$ Cholesterol degradation rate: We utilise the value previously derived ⁹⁹⁶ in Bhattacharya et al. [2] of 1.20×10^{-4} /s.
- ⁹⁹⁷ $\bar{\omega}$ **HMGCoA production rate**: This value has been determined from our ⁹⁹⁸ sensitivity analysis to be 3.895×10^{11} molec./ml. The value has been found to ⁹⁹⁹ ensure enough cholesterol is produced.
- 1000 $\mathbf{x_h}$ Number of binding sites for SREBP-2 on HMGCR DNA: Vallett 1001 et al. [28] state a value of 3.
- $\mathbf{x_s}$ Number of binding sites for SREBP-2 on squalene synthase **DNA**: Without further evidence we assume this is 1.
- $\mathbf{x_{c}}$ Number of binding sites on SREBP-2 for cholesterol: Radhakrishnan et al. [46, 16] state a value of 4.

1006 References

- [1] August, E., Parker, K., Barahona, M., 2007. A dynamical model of
 lipoprotein metabolism. Bulletin of Mathematical Biology 69 (4), 1233–
 1254.
- [2] Bhattacharya, B., Sweby, P., Minihane, A.-M., Jackson, K., Tindall,
 M., 2014. A mathematical model of the sterol regulatory element binding protein 2 cholesterol biosynthesis pathway. Journal of Theoretical
 Biology 349, 150–162.
- [3] Brown, M., Dana, S., Goldstein, J., 1974. Regulation of 3-hydroxy-3methylglutaryl coenzyme A reductase activity in cultured human fibroblasts comparison of cells from a normal subject and from a patient with homozygous familial hypercholesterolemia. Journal of Biological Chemistry 249 (3), 789–796.
- [4] Brown, M., Goldstein, J., 1980. Multivalent feedback regulation of HMG
 CoA reductase, a control mechanism coordinating isoprenoid synthesis
 and cell growth. Journal of Lipid Research 21 (5), 505–517.

- In 1022 [5] Bruenger, E., Rilling, H., 1988. Determination of isopentenyl diphosphate and farnesyl diphosphate in tissue samples with a comment on secondary regulation of polyisoprenoid biosynthesis. Analytical Biochemistry 173 (2), 321–327.
- [6] Buhaescu, I., Izzedine, H., 2007. Mevalonate pathway: a review of clinical and therapeutical implications. Clinical Biochemistry 40 (9), 575–
 584.
- ¹⁰²⁹ [7] Cole, S., Vassar, R., 2006. Isoprenoids and alzheimer's disease: A com-¹⁰³⁰ plex relationship. Neurobiology of Disease 22, 209–222.
- [8] Darzacq, X., Shav-Tal, Y., de Turris, V., Brody, Y., Shenoy, S., Phair,
 R., Singer, R., 2007. In vivo dynamics of RNA polymerase II transcription. Nature Structural & Molecular Biology 14 (9), 796–806.
- [9] DeBose-Boyd, R., 2008. Feedback regulation of cholesterol synthesis:
 sterol-accelerated ubiquitination and degradation of hmg coa reductase.
 Cell Research 18, 609–21.
- [10] Foresti, O., Ruggiano, A., Hannibal-Bach, H., Ejsing, C., Carvalho,
 P., 2013. Sterol homeostasis requires regulated degradation of squalene
 monooxygenase by the ubiquitin ligase Doa10/Teb4. Elife 2, e00953.
- [11] Gaylor, J. L., 2002. Membrane-bound enzymes of cholesterol synthesis
 from lanosterol. Biochemical and biophysical research communications
 292 (5), 1139–1146.
- [12] Goldstein, J., Brown, M., 1984. Progress in understanding the LDL
 receptor and HMG-CoA reductase, two membrane proteins that regulate
 the plasma cholesterol. Journal of Lipid Research 25 (13), 1450–1461.
- ¹⁰⁴⁶ [13] Goldstein, J., Brown, M., 1990. Regulation of the mevalonate pathway. ¹⁰⁴⁷ Nature 343 (6257), 425–30.
- [14] Hampton, R., 2002. Proteolysis and sterol regulation. Annual Review of
 Cell and Developmental Biology 18, 345–78.
- [15] Istvan, E., Palnitkar, M., Buchanan, S., Deisenhofer, J., 2000. Crystal structure of the catalytic portion of human HMG-CoA reductase:
 insights into regulation of activity and catalysis. The EMBO Journal
 19 (5), 819–830.

- [16] Jackson, K., Maitin, V., Leake, D., Yaqoob, P., Williams, C., 2006.
 Saturated fat-induced changes in Sf 60–400 particle composition reduces
 uptake of LDL by HepG2 cells. Journal of Lipid Research 47 (2), 393–403.
- [17] Jiang, G., McKenzie, T., Conrad, D., Shechter, I., 1993. Transcriptional regulation by lovastatin and 25-hydroxycholesterol in HepG2 cells and molecular cloning and expression of the cDNA for the human hepatic squalene synthase. Journal of Biological Chemistry 268 (17), 12818– 12824.
- [18] Kellner-Weibel, G., Geng, Y., Rothblat, G., 1999. Cytotoxic cholesterol
 is generated by the hydrolysis of cytoplasmic cholesteryl ester and transported to the plasma membrane. Atherosclerosis 146, 309–19.
- [19] Kervizic, G., Corcos, L., 2008. Dynamical modeling of the cholesterol regulatory pathway with boolean networks. BMC Systems Biology
 2 (99), doi: 10.1186/1752-0509-2-99.
- [20] Lickwar, C. R., Mueller, F., Hanlon, S. E., McNally, J. G., Lieb, J. D.,
 2012. Genome-wide protein-DNA binding dynamics suggest a molecular
 clutch for transcription factor function. Nature 484 (7393), 251–255.
- 1072 [21] Matlab, Release 2016b. The MathWorks. Natick, Massachusetts, United
 1073 States.
- [22] Mazein, A., Watterson, S., Hsieh, W., WJ, G., P, G., 2013. A comprehensive machine-readable view of the mammalian cholesterol biosynthesis pathway. Biochemical Pharmacology 86 (1), 56–66.
- [23] Menegola, E., Broccia, M., Di Renzo, F., E, G., 2006. Postulated
 pathogenic pathway in triazole fungicide induced dysmorphogenic effects. Reproductive Toxicology 260 (2), 186–95.
- [24] Michaelis, L., Menten, M., 1913. Die kinetik der invertinwirkung. Biochemische Zeitschrift 49 (333-369), 352.
- [25] Morgan, A., Mooney, K., Wilkinson, S., Pickles, N., MT., M. A., 2016.
 Mathematically modelling the dynamics of cholesterol metabolism and ageing. Biosystems 145, 19–32.

- [26] Murphy, L., Moore, T., S, N., 2012. Propiconazole-enhanced hepatic cell
 proliferation is associated with dysregulation of the cholesterol biosynthesis pathway leading to activation of Erk1/2 through Ras farnesylation. Toxicology and Applied Pharmacology 260 (2), 146–54.
- ¹⁰⁸⁹ [27] Murray, J., 2002. Mathematical Biology I: An Introduction. Springer,
 ¹⁰⁹⁰ New York, USA.
- [28] Radhakrishnan, A., Goldstein, J., McDonald, J., Brown, M., 2008.
 Switch-like control of SREBP-2 transport triggered by small changes in ER cholesterol: a delicate balance. Cell Metabolism 8 (6), 512–521.
- [29] Radhakrishnan, A., Sun, L.-P., Kwon, H., Brown, M., Goldstein, J.,
 2004. Direct binding of cholesterol to the purified membrane region of
 SCAP: mechanism for a sterol-sensing domain. Molecular Cell 15 (2),
 259–268.
- [30] Rudling, M., Angelin, B., Ståhle, L., Reihnér, E., Sahlin, S., Olivecrona, H., Björkhem, I., Einarsson, C., 2002. Regulation of hepatic low-density lipoprotein receptor, 3-hydroxy-3-methylglutaryl coenzyme A reductase, and cholesterol 7α -hydroxylase mRNAs in human liver. The Journal of Clinical Endocrinology & Metabolism 87 (9), 4307–4313.
- [31] Sanguinetti, G., Lawrence, N., Rattray, M., 2006. Probabilistic infer ence of transcription factor concentrations and gene-specific regulatory
 activities. Bioinformatics 22 (22), 2775–2781.
- [32] Sebti, S., Hamilton, A., 1997. Inhibition of Ras prenylation: A novel approach to cancer chemotherapy. Pharmacology and Therapeutics 74 (1), 103–114.
- [33] Segal, I., 1975. Enzyme kinetics behaviour and analysis of rapid equilibrium and steady-state enzyme systems. A Wiley-Interscience Publication, New York, USA.
- [34] Siperstein, M., Fagan, V., 1966. Feedback control of mevalonate synthesis by dietary cholesterol. Journal of Biological Chemistry 241 (3),
 602–609.
- [35] Srivastava, R., Ito, H., Hess, M., Srivastava, N., Schonfeld, G., 1995.
 Regulation of low density lipoprotein receptor gene expression in HepG2

- and Caco2 cells by palmitate, oleate, and 25-hydroxycholesterol. Journal of lipid research 36 (7), 1434–1446.
- [36] Tanaka, R., Edwards, P., Lan, S., Knöppel, E., Fogelman, A., 1982.
 Purification of 3-hydroxy-3-methylglutaryl coenzyme A reductase from human liver. Journal of Lipid Research 23 (4), 523–530.
- [37] Tansey, T., Shechter, I., 2000. Structure and regulation of mammalian
 squalene synthase. Biochimica et Biophysica Acta-Molecular and Cell
 Biology of Lipids 1529 (1), 49–62.
- ¹¹²⁵ [38] The Metabolomics Innovation Centre, Accessed: 31-07-2015. The
 ¹¹²⁶ human metabolome database. http://www.hmdb.ca/metabolites/
 ¹¹²⁷ HMDB01375.
- ¹¹²⁸ [39] Trachsel, H., 1991. Translation in eukaryotes. CRC Press, Florida, USA.
- [40] Turley, S., Dietschy, J., 2003. The intestinal absorption of biliary and dietary cholesterol as a drug target for lowering the plasma cholesterol level. Preventive Cardiology 6 (1), 29–64.
- [41] Cards, G., Accessed: 15-10-2017. Hmgcr gene size. http://www.
 genecards.org/cgi-bin/carddisp.pl?gene=HMGCR.
- [42] DNA technologies, I., Accessed: 17-10-2017. Molecular facts and figures.
 https://www.idtdna.com/pages/docs/educational-resources/
 molecular-facts-and-figures.pdf?sfvrsn=4.
- [43] van Greevenbroek, M., de Bruin, T., 1998. Chylomicron synthesis by
 intestinal cells in vitro and in vivo. Atherosclerosis 141, S9–S16.
- [44] Vargas, N., Brewer, B., Rogers, T., Wilson, G., 2009. Protein kinase
 C activation stabilizes LDL receptor mRNA via the JNK pathway in
 HepG2 cells. Journal of Lipid Research 50 (3), 386–397.
- [45] Watterson, S., Guerriero, M., Blanc, M., Mazein, A., Loewe, L., Robertson, K., Gibbs, H., Shui, G., Wenk, M., Hillston, J., P., G., 2013. A
 model of flux regulation in the cholesterol biosynthesis pathway: Immune mediated graduated flux reduction versus statin-like led stepped
 flux reduction. Biochimie 95 (3), 613–21.

- [46] Wilson, G., Deeley, R., 1995. An episomal expression vector system for
 monitoring sequence-specific effects on mRNA stability in human cell
 lines. Plasmid 33 (3), 198–207.
- [47] Yang, W., Swartz, J., 2011. A filter microplate assay for quantitative analysis of DNA binding proteins using fluorescent DNA. Analytical Biochemistry 415 (2), 168–174.