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Mathematically Modelling the Dynamics of Cholesterol Metabolism and Ageing

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Keywords

Cholesterol, ageing, low density lipoprotein cholesterol (LDL-C), high density lipoprotein cholesterol (HDL-C), cholesterol ester transfer protein (CETP)

Abstract

Cardiovascular disease (CVD) is the leading cause of morbidity and mortality in the UK. This condition becomes increasingly prevalent during ageing; 34.1% and 29.8% of males and females respectively, over 75 years of age have an underlying cardiovascular problem. The dysregulation of cholesterol metabolism is inextricably correlated with cardiovascular health and for this reason low density lipoprotein cholesterol (LDL-C) and high density lipoprotein cholesterol (HDL-C) are routinely used as biomarkers of cardiovascular disease (CVD) risk. The aim of this work was to use mathematical modelling to explore how cholesterol metabolism is affected by the ageing process. To do this we updated a previously published whole-body mathematical model of cholesterol metabolism to include an additional 96 mechanisms that are fundamental to this biological system. Additional mechanisms were added to cholesterol absorption, cholesterol synthesis, reverse cholesterol transport (RCT), bile acid synthesis, and their enterohepatic circulation. The sensitivity of the model was explored by the use of both local and global parameter scans. In addition, acute cholesterol feeding was used to explore the effectiveness of the regulatory mechanisms which are responsible for maintaining whole-body cholesterol balance. It was found that our model behaves as a hypo-responder to cholesterol feeding, while both the hepatic and intestinal pools of cholesterol increased significantly. The model was also used to explore the effects of ageing in tandem with three different CETP genotypes. Ageing in the presence of an atheroprotective CETP genotype, conferring low CETP activity, resulted in a 0.6% increase in LDL-C. In comparison, ageing with a genotype reflective of high CETP activity, resulted in a 1.6% increase in LDL-C. Thus, the model has illustrated the importance of the I405V CETP genotype and its potential role in healthy ageing.

1.0. Introduction

Life expectancy is increasing at a phenomenal rate. If one examines life expectancy in the UK at the beginning of the 1980s and compares it to the projected value for 2081, then this demographic shift in favour of older people is truly emphasized. Males and females born in 1981, had a life expectancy of 70.9 and 76.9 years respectively, while the projected values for 2081 are 89.7 and 92.5 years (OFNS, 2013). It is also anticipated that the percentage of individuals in the UK over 60 years will double to 22% by 2050, when compared to 2000 (WHO, 2014). Moreover by 2050 it is anticipated, that the number of individuals >80 years will quadruple (WHO, 2014). This increase in life expectancy comes with significant challenges for older people. Most significant of these is maintaining health status by remaining disease free. Of the diseases which burden older people in the UK, CVD is the leading cause of morbidity (30.3%) in individuals ≥ 60 years of age (Prince et al., 2015). CVD has many risk factors (Appelman et al., 2015), however the plasma levels of total cholesterol, LDL-C and HDL-C are long established gold standard risk factors for CVD (Appelman et al., 2015). Specifically, rises in both total and LDL-C result in an increase in CVD risk, while conversely a rise in HDL-C results in a decrease in CVD risk (Austin et al., 1988; Cooney et al., 2009). On a population basis, ageing has been shown to result in a rise in both total and LDL-C in males and females (Abbott et al., 1983). For example, data from the Framingham Study demonstrates there is a gradual rise in LDL-C from an initial value of ~ 100 mg/dL in 15-19 year olds (males and females), to 143 and 159mg/dL in 60-64 year olds, males and females respectively. Beyond the 6th decade, LDL-C levels continue to rise in females, however they marginally decline in males within this population (Abbott et al., 1983). The most likely reason for this is that ageing results in the dysregulation of whole-body cholesterol metabolism (Mooney and Mc Auley, 2015). Whole-body cholesterol metabolism is regulated by a number of factors including cholesterol synthesis, intestinal cholesterol absorption, hepatic cholesterol uptake (known as reverse cholesterol transport), cholesterol excretion, bile acid production, and deconjugation by intestinal microflora and subsequent excretion. Changes to any of these mechanisms can dysregulate cholesterol metabolism (Morgan et al., 2016).

The multifaceted nature of cholesterol metabolism, the complexities of its dynamics, and the long term impact of the ageing process on its behaviour makes studying this system a challenge (Mc Auley and Mooney, 2014). In recent years however, several groups have adopted a systems level approach to investigating lipid metabolism (Mc Auley and Mooney, 2015; Mooney et al., 2016). The majority of these models have centred on specific metabolic processes, including those directed at lipoprotein metabolism (Hubner et al., 2008; Shorten and Upreti, 2005; Sips et al., 2014), cholesterol biosynthesis (Bhattacharya et al., 2014; Kervizic and Corcos, 2008; Mazein et al., 2013; Watterson et al., 2013), reverse cholesterol transport (Lu et al., 2014), adipocyte metabolism (Micheloni et al., 2015), hepatocyte metabolism (Jerby et al., 2010), cholesterol regulatory enzymes (Chapman et al., 2010), whole-body plasma cholesterol metabolism (van de Pas et al., 2012) and enterohepatic circulation of bile acids (Mishra et al., 2014). These models all have noteworthy features and have added to our understanding of lipid metabolism. Among the computational systems biology models, the model by Mc Auley et al (2012) provides an integrated template for investigating ageing and whole-body cholesterol metabolism (Mc Auley et al., 2012). This model is encoded in the systems biology markup language (SBML) (Hucka et al., 2004; Hucka et al., 2003), and is part of the curated section in the BioModels web site; a repository for models encased in SBML exchange framework (Le Novère et al., 2006; Li et al., 2010). This makes the model straightforward to adapt and update. The adaptable nature of this model was emphasized recently by Mishra, Somvanshi & Venkatesh (2014), who used this model in tandem with another model (Demirezen and Barlas, 2008). Their combined model included the variables body weight, diet and exercise and they analysed the feedback mechanisms of enterohepatic circulation of bile acids; specifically, bile acid synthesis, interactions with dietary lipids, and excretion. Despite its integrated nature and its focus on ageing, the model by Mc Auley et al. (2012) is lacking several key mechanisms that are fundamental to cholesterol metabolism. These key mechanisms include plasma membrane receptors, nuclear receptors and a variety of *in vivo* and intestinal microfloral enzymes. It is imperative that these factors are included in any holistic representation of cholesterol metabolism. Thus, the aims of this work were fourfold. 1) To mechanistically update the whole-body mathematical model of cholesterol metabolism by Mc Auley et al. (2012). 2) To use the

updated model to explore the role of cholesterol and saturated fat (SFA) feeding on lipoprotein profile. 3) To examine the role of age-related alterations to cholesterol metabolism on the lipoprotein profile, specifically, cholesterol absorption, LDL-C clearance, and bile acid synthesis and subsequent deconjugation. 4) To test the model using pharmacological interventions that have been used *in vivo* to treat hypercholesterolaemia.

2.0. Overview of Cholesterol Metabolism

Maintaining cholesterol metabolism is a subtle balancing act between the amount of cholesterol ingested, absorbed, excreted and synthesised (Mc Auley et al., 2005). Ingested cholesterol, exists as esters, and free cholesterol (Iqbal and Hussain, 2009). Esterified cholesterol is hydrolysed to free cholesterol by cholesterol ester hydrolase (CEH)/cholesterol esterase/bile salt-dependent lipase (BSDL) for micelle incorporation (Ikeda et al., 2002). This facilitates the absorption into the jejuncyte by Neimann-Pick C1-Like 1 protein (NPC1L1) endocytosis (Jia et al., 2011; Petersen et al., 2008). Internalized free cholesterol is converted to esterified cholesterol by acyl coenzyme A: cholesterol acyltransferase 2 (ACAT2). It is then transported to apolipoprotein (apo) B-48 to form part of a nascent chylomicron via microsomal triglyceride transfer protein (MTP) (Atzel and Wetterau, 1993; Atzel and Wetterau, 1994). Acquisition of apo C-II and E converts the chylomicron to a mature chylomicron. On delivery to adipose/muscle tissue, apo C-II activates of lipoprotein lipase (LPL), which hydrolysis TAG releasing glycerol and FFAs (Kersten, 2014; Olivecrona and Beisiegel, 1997). Hepatocytes then uptake the chylomicron remnants via LDLr and low density lipoprotein receptor-related protein (LRP), for degradation (Cooper, 1997). Reverse cholesterol transport (RCT) removes excess cholesterol from peripheral tissues and transports it to the liver for excretion in bile or conversion to bile acids (Ohashi et al., 2005; Scheibner et al., 1994). RCT begins with the efflux of cholesterol to nascent HDLs (ndHDL) (Lund-Katz et al., 2013). Cholesterol efflux can occur via the receptors ABC-A1 (He et al., 2015) or scavenger receptor class B type 1 (SR-B1) (Ji et al., 2011), or via passive diffusion (Gillotte et al., 1998). Lecithin-cholesterol acyltransferase (LCAT) esterifies the incorporated cholesterol, converting the ndHDL to HDL₃ (Sorci-Thomas et al., 1990). The assembly of 2 small, dense, HDL₃ molecules, in the presence of phospholipid transfer protein (PLTP), induces the formation of a larger HDL₂ molecule (Chirackal Manavalan et al., 2014), which can be reversed by hepatic lipase (HL) (Patsch et al., 1987). Cholesterol esters can enter the liver directly via the receptor SR-B1, or via CETP, which facilitates the 1:1 exchange of cholesterol esters from HDL₂ with TAGs from very low density lipoprotein (VLDL) and LDL (Zhang et al., 2015). These lipoproteins return the esterified cholesterol to the liver (Brown and Goldstein, 1984). LDL can also return cholesterol esters to peripheral tissues (Baigent et al., 2010). VLDLs and IDLs that haven't deposited cholesterol esters into the liver, can be metabolised via LPL and hormone sensitive lipase (HSL) to LDL.

The enterohepatic circulation of bile acids occurs between 4-12 times per day, but only 5% per day are lost in faeces (Gerard, 2013), and of this, 98% is in the unconjugated form (Batta et al., 1999). Although preferentially hydrolysing glyco-conjugated bile acids, both glyco- and tauro-conjugated bile acids can be converted to unconjugated bile acids whilst in the intestinal lumen by the hydrolysis of the amine bond, and removal of glycine or taurine by bacterial bile salt hydrolase (BSH) (Jarocki et al., 2014). Certain intestinal microflora have the ability to produce BSH, but *Lactobacillus* species are responsible for 86% of the BSH activity in the ileum, and 74% in the caecum (Tannock et al., 1989). The probiotic administration of BSH-producing strains can lead to a decrease of 63.2% in plasma LDL-C levels. This is due to the increase in excreted bile acids, leading to enhanced *de novo* synthesis of bile acids from cholesterol, to replace those excreted. Furthermore, there is a decrease in cholesterol solubility due to an increase in the proportion of unconjugated bile acids. In addition, there is a rise in the ratio of propionate to acetate (Begley et al., 2006; Guo and Li, 2013). However, this treatment strategy is controversial as evidence suggests that the byproducts of unconjugated bile acids have been associated with the dysregulation of mucosal functioning (Ajouz et al., 2014; Baptissart et al., 2013).

3.0. How Cholesterol Metabolism changes with Age

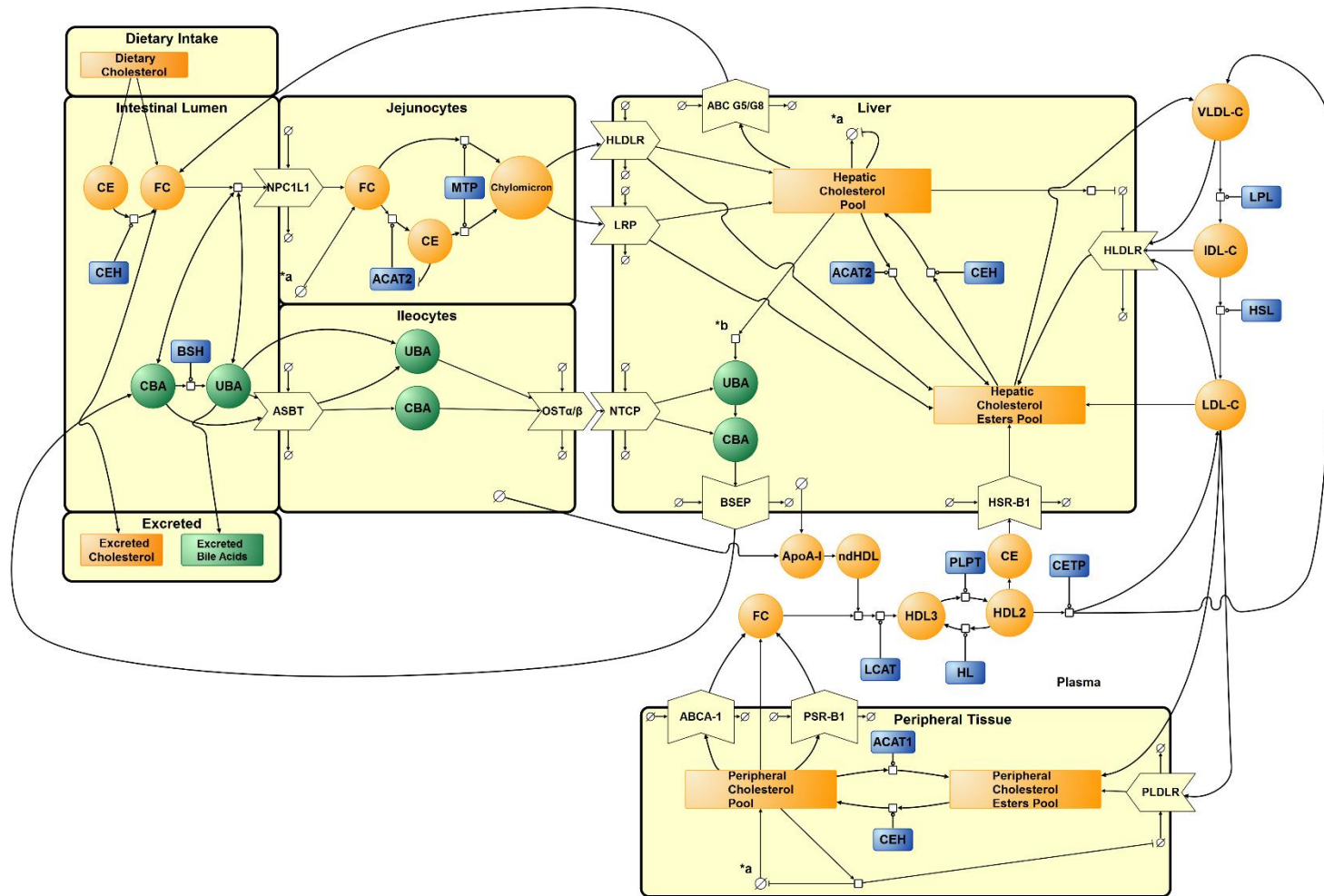
It has been consistently shown that LDL-C increases with age in both males and females across different population groups (Abbott et al., 1983; Yang et al., 2012). A number of mechanisms have been implicated in this age-related rise in LDL-C. For example, it has been suggested that an increase in the absorption of dietary cholesterol is one of the factors responsible for the increase of serum LDL-C with age (Duan et al., 2006; Hollander and Morgan, 1979). Murine models demonstrate that this increase in cholesterol absorption is correlated with the age-related increase in the apical membrane receptor NPC1L1 in the duodenum and jejunum (Duan et al., 2006; Hollander and Morgan, 1979). Ageing resulted in a 17% increase in cholesterol absorption in one murine model due to a significant increase in NPC1L1 mRNA in both the duodenum and jejunum (Duan et al., 2006). The receptor NPC1L1 plays a vital role in the absorption of dietary cholesterol. Thus, inhibiting NPC1L1 with ezetimibe is one strategy that has been adopted in the treatment of hypercholesterolaemia (Dujovne et al., 2002). Another mechanism that has been associated with the dysregulation of cholesterol metabolism is a decrease in LDLr numbers and/or activity. LDLr is responsible for the uptake of circulating LDL-C from the plasma. LDLr therefore lowers the plasma LDL-C levels and reduces CVD risk (Brown and Goldstein, 1984). The age-related decline in LDLr, results in a reduction in LDL clearance, thus raising the plasma LDL-C levels (Millar et al., 1995). This decline in LDLr with age could be due to the age-related increase in proprotein convertase subtilisin kexin-9 (PCSK9), a serine protease associated with LDLr degradation (Cui et al., 2010). Additionally, this could be due to the decreased demand for cholesterol for the *de novo* synthesis of bile acid, due to the age-related decrease in bile acid synthesis (Ericsson et al., 1991). A decline in bile acid synthesis and hepatic expression of CYP7A1 have been shown to be correlated with ageing in human subjects (Bertolotti et al., 2007). This decreases cholesterol utilisation for the *de novo* synthesis of bile acids (Bertolotti et al., 2007). This is supported by Bertolotti et al. (1993) who carried out a linear regression analysis which estimated that the amount of cholesterol undergoing the CYP7A1 regulated reaction in humans decreases by 60mg per day, every 10 years (Bertolotti et al., 1993). It is also possible that the down regulation of CYP7A1 is due to the age related decline of growth hormone (GH), as illustrated by rodent studies (Parini et al., 1999). Similar studies have also illustrated that sex hormones play a role in the decline of CYP7A1, as gender differences are often observed. For example, aged female rodents often have an increased concentration of serum bile acids, whereas levels remain constant in males with age (Fu et al., 2012). This is in addition to alterations to the composition of bile acids (Trautwein et al., 1999). Gut microflora are also affected by the ageing process. Levels and species diversity of *Lactobacillus* and *Bifidobacterium* species are consistently lower in older people than younger human adults (Hopkins and Macfarlane, 2002; Hopkins et al., 2001). These species are well documented for their ability to produce the enzyme BSH, which is responsible for the deconjugation of bile acids in the small intestine (Tanaka et al., 1999). This decrease in BSH-positive species leads to a decrease in bile acid excretion and thus a decrease in cholesterol utilisation for the *de novo* synthesis of replacement bile acids (Uchida et al., 1996). This could be one cause of the increase in LDL-C with age. Interestingly, supplementation with probiotic strains in elderly rodent populations has been shown to reduce LDL-C levels (Al-Sheraji et al., 2012) as well as improve gut health (Tuohy et al., 2003) and increase immunity in human subjects (Moro-Garcia et al., 2013). Mutations in the gene coding for CETP have also been linked to healthy ageing. CETP mediates the transport of cholesterol esters from HDL to LDL and VLDL. Mutation to the CETP gene has been associated with longevity in Ashkenazi Jews (Atzmon et al., 2005; Barzilai et al., 2003). The amino acid substitution of 405 isoleucine to valine (I405V) in the CETP gene results in a lower serum CETP concentration and subsequently significantly larger LDL and HDL. The CETP I405V genotype is associated with a reduced prevalence of CVD, diabetes mellitus, and hypertension, possibly contributing to the exceptional longevity observed in these individuals (Atzmon et al., 2005; Barzilai et al., 2003). The assertion that reduced CETP levels convey a protective phenotype for healthy ageing, has led to investigations into the use of CETP inhibitors as a therapeutic strategy for the treatment of hypercholesterolaemia. For example, it has been recently revealed that in phase II clinical trials the CETP inhibitor TA-8995 decreased LDL-C by 27.4% with a 1mg daily dose, with minimal adverse effects (Hovingh et al., 2015). Therefore, with our updated model, there was a clear emphasis on these aspects of lipid metabolism.

4.0. Methods

4.1. Diagrammatic Representations of Whole-Body Cholesterol Metabolism

Our updated model consists of 8 compartments which are: 1) dietary intake, 2) intestinal lumen, 3) jejunocytes, 4) ileocytes, 5) hepatic tissue, 6) blood, 7) peripheral tissue and 8) excreted (**Figure 1**). Figures 1 – 3, were created using systems biology graphical notation (SBGN). SBGN is a graphical representation modelling format designed to be employed as a universal framework for the construction of diagrammatic representations of biological or other processes, to allow easy recognition of each species and process (Le Novere et al., 2009). Figure 1 is an adaptation of the previous SBGN diagram generated by Mc Auley et al. (2012). The SBGN diagrams were developed with VANTED (Version 2.2.1, <http://vanted.ipk-gatersleben.de/>, http://www.sbgng.org/Main_Page) (Rohn et al., 2012). To incorporate the additional information into the SBGN diagram, the compartment previously referred to as 'intestine' was divided into 3 sub-compartments, 'intestinal lumen', 'jejunocytes' and 'ileocytes'. The role of several key processes, receptors and enzymes could then be incorporated. These include the differentiation of dietary free and esterified cholesterol, bile acid micelle formation for the uptake of free cholesterol via the NPC1L1 receptor, cholesterol esterification in the jejunum, chylomicron transport, and bile acid reabsorption. The mechanisms underpinning RCT were expanded from those represented in the original model. The enzymes PLPT, HL, LCAT and CETP and their roles in HDL subtype composition and cholesterol transport were included. Following this, additional receptors were incorporated into the peripheral tissue, hepatic tissue, and the newly formed ileocytes compartments to further elucidate the transport mechanisms of cholesterol, lipoproteins and bile acids. Next, cholesterol biosynthesis was converted from a 1 step process in the Mc Auley et al. (2012) model to an 18 step process. Overall, cholesterol biosynthesis has significantly more steps than this. However, the rationale for abstracting to an 18 step reaction system is as follows. The first 13 steps of cholesterol biosynthesis are well characterised (Liscum, 2008). The latter reactions in this process remain poorly characterised. Therefore, the reactions that were included are those that are central to the pathway and are well delineated. Due to the size and complexity of this reaction network, a separate SBGN diagram was developed to illustrate this component of the model (**Figure 2**). In our overall diagrammatic representation of the model (Figure 1) these reactions are simplified to 3 source and sink glyphs labelled *a, in the jejunocytes, hepatic and peripheral tissue compartments. Finally, the *de novo* synthesis of bile acids from hepatic cholesterol was converted from a 1 step process to a 15 step process, and similar to the cholesterol synthesis pathway, was too detailed to be represented fully in Figure 1. Bile acid synthesis is therefore represented by the source and sink glyph labelled *b in Figure 1, and is fully illustrated in **Figure 3**. The reactions outlined in Figure 3 are derived from various sources. In essence they are a summary of the processes that are well characterised. Some well characterised steps are excluded from the summary to reduce the process diagram size, and for clarity. For example, following the chain shortening of 4-cholesten-7 α , 12 α - diol-3-one by CYP27A1 further modification by SLC27A5 (EC 6.2.1.7), ACOX2 (EC 1.17.99.3), HSD1724 (EC 1.1.135) and SCP2 (EC 2.3.1.176), occurs before cholic acid formation. There is ambiguity surrounding the re-entry of compounds involved in the alternative pathways to the final steps before the formation of cholic or deoxycholic acid resulted in gaps of knowledge. For this reason, there are process glyphs that represent a series of uncertain steps.

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20 Figure 1. SBGN diagram of mathematical model of whole-body cholesterol metabolism. The diagram details the transport of cholesterol, bile acids
 21 and lipoproteins through cholesterol absorption, lipoprotein metabolism, RCT and enterohepatic circulation. For abbreviations see Table A.2. Round
 22 arrow head represent the target of a catalytic enzyme, represented as blue rounded rectangles. Arrows represent flux. Hatched process nodes represent omitted
 23 processes. T shaped arrows represent inhibition. The Greek symbol theta represents synthesis/degradation. a* cholesterol synthesis (see Figure 2),
 24 b* bile acid synthesis (see Figure 3).

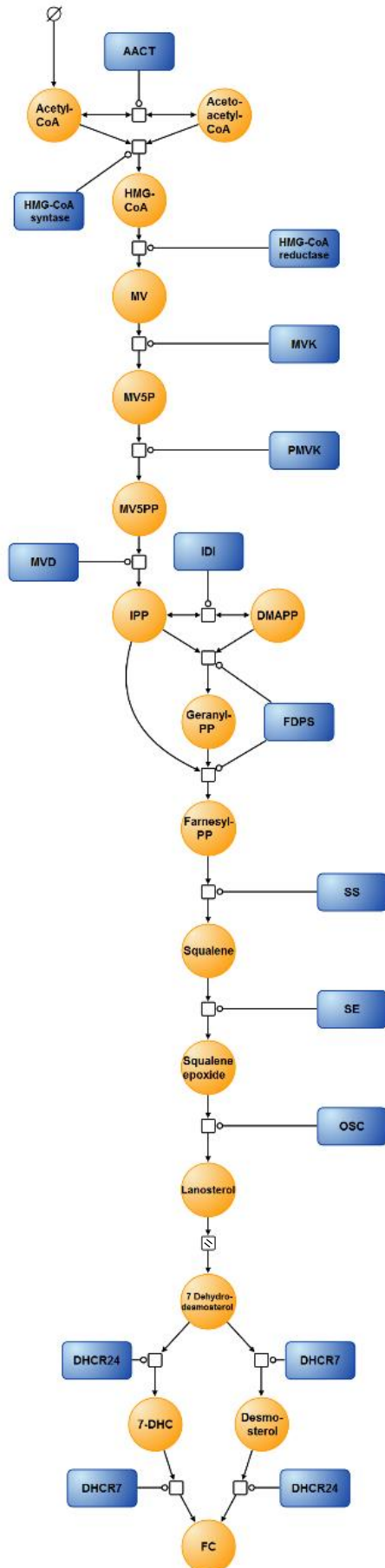


Figure 2. Cholesterol biosynthesis. Synthesis of cholesterol from acetyl CoA via intermediates HMG CoA, mevalonate, squalene and lanosterol. See Table A.2 for abbreviations. Round arrow heads represent the target of a catalytic enzyme, represented as blue rounded rectangles. Arrows represent flux. Hatched process nodes represent omitted processes. The Greek symbol theta represents synthesis. Acetoacetyl CoA thiolase catalyses the interconversion of acetyl CoA and acetoacetyl CoA. Acetyl CoA and acetoacetyl CoA undergo a condensation reaction to form 3-hydroxy-3-methylglutaryl-coenzyme A (HMG CoA). HMG CoA is then converted by HMG CoA reductase to mevalonate. Phosphorylation of mevalonate forms mevalonate-5P, which is further phosphorylated to mevalonate-5PP. Decarboxylation and dehydration of mevalonate-5PP forms isopentenyl-PP (IPP), which converts to its isoform dimethylallyl-PP (DMAPP). DMAPP reacts with IPP to create geranyl-PP. Further condensation and the addition of another IPP creates farnesyl-PP. Condensation of 2 farnesyl-PP molecules forms squalene, which is converted to squalene epoxide before undergoing cyclisation to form lanosterol. A series of reactions, including the branching of 7-dehydrodesmosterol to either desmosterol or 7-dehydrocholesterol, both of which can then be converted to cholesterol.

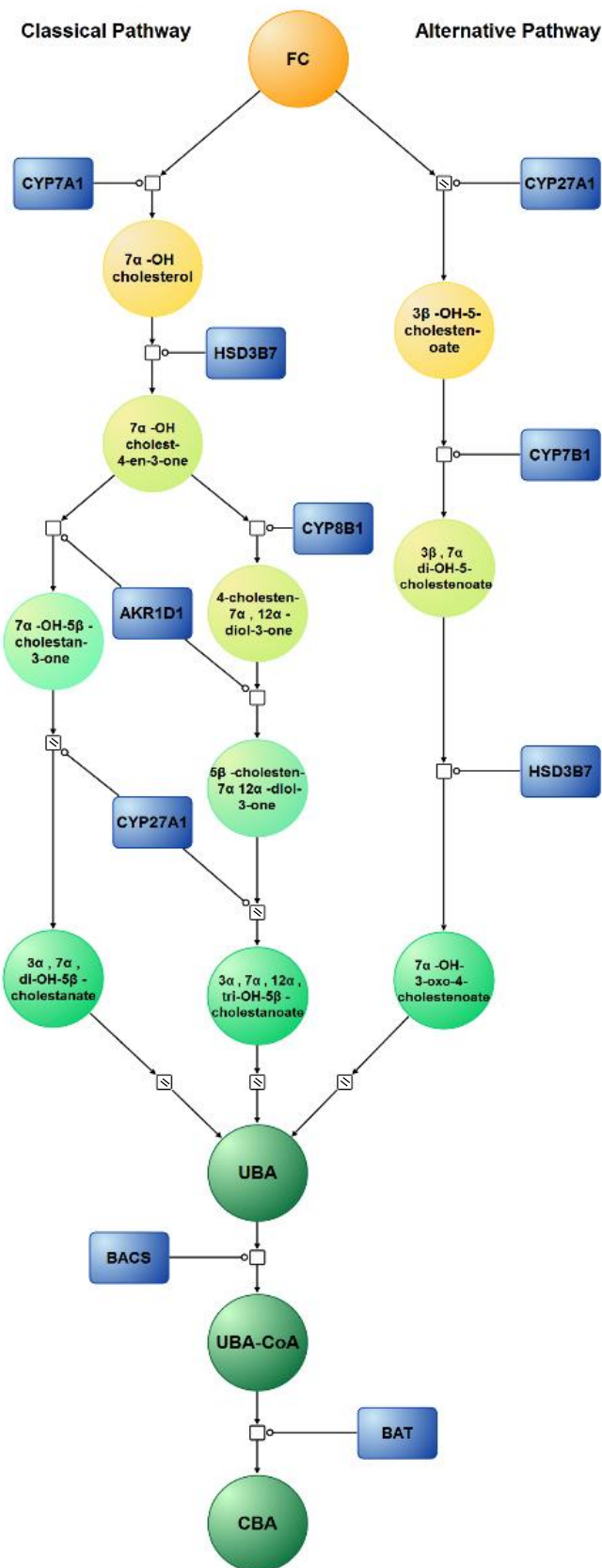


Figure 3. Bile Acid Synthesis. Classical Pathway: Hydroxylation of cholesterol creates 7 α -hydroxycholesterol. Subsequent modification of the sterol ring forms 7 α -hydroxy-5 β -cholesten-3-one before the classical pathway splits, to enable the formation of either cholic acid (CA) or chenodeoxycholic acid (CDCA). For the formation of cholic acid, modification of the sterol ring by 12 α -hydroxylase (CYP7B1) must occur before further modification. The side chain of 5 β -cholesten-7 α , 12 α -diol-3-one is then acted upon by a series of oxidation steps leading to its shortening. Further modification by several enzymes then takes place to form CA. The same steps, although in the absence of CYP8B1, are undergone in the other route of the classical pathway, to form CDCA. Alternative Pathway: Side chain shortening occurs first in the alternative pathway creating 27 α -hydroxycholesterol, before hydroxylation creates 3 β -7 α -dihydroxy-5-cholesten-3-one. HSD3B7, CYP8B1 and AKR1D1 modification of the sterol ring can then occur, and upon further alteration, CA and CDCA can then be formed. This is followed by conjugation of 98% of these bile acids. They are effluxed from the hepatocyte primarily by bile salt export pump (BSEP) in the rate limiting step of bile acid circulation. See Table A.2 for abbreviations. Round arrow head represent the target of catalytic enzyme, represented as blue rounded rectangles. Arrows represent flux. Hatched process nodes represent omitted processes.

4.2. Model Assembly and Parameterisation

The SBML for the original model was downloaded from the BioModels database (Juty et al., 2015) (<https://www.ebi.ac.uk/biomodels-main/>, BIOMD0000000434). It was then imported into Version 4.14.89 of Copasi (Hoops et al., 2006). Within Copasi the model was then reassembled. The original model contained 48 species. The updated version now contains 144 species including the 4 species used as counters for the event reactions (**Table A.3**). The first processes to be added were the ingestion of free cholesterol and cholesterol esters. To represent these reactions we defined a number of new species, and parameters and then introduced two SBML events for each species. An event in SBML is a reaction which is activated at a particular point in time and which causes a change in the value of a species concentration, compartment size, or parameter. In this case, the rationale for including events was to represent the ingestion of cholesterol every 8 hours. This was more biologically realistic than the single reaction which represented this process previously. The events were defined in using the syntax outlined in **Table A.1**. Events stimulated the intake of both free cholesterol and cholesterol esters. The creation of the events reflected the intake of a meal every 8 hours. Following the assembly of the events, the entry of dietary esterified cholesterol into the intestinal lumen 3 times a day was deterministically simulated (**Figure 4A**). Once in the intestinal lumen, esterified cholesterol is converted to free cholesterol for entry into the jejuncytes. **Figure 4B** illustrates how the level of free cholesterol in the jejuncyte was affected by the entry of this converted free cholesterol. This is in addition to the 88.667mg of free cholesterol that is also ingested 3 times a day, and the 1000mg/day of biliary cholesterol effluxed back into the lumen from the liver. Within the model, free cholesterol contained in the jejunum fluctuated. Initial steady state values for free and esterified cholesterol in the hepatic or peripheral tissues were calculated based on the experimental data in **Table 1** (**Figure 4C**). Other species were calculated in a similar way, utilising tissue mass and molecular mass. The initial steady state levels of many other intermediate species were assumed. Each value is in mg/compartment. **Table A.2** provides the details of the species and their abbreviations, while **Table A.3** provides details of over 144 kinetic based reactions which in total make up the new version of the model. To summarise the kinetics, the model includes 37 reactions which are derived using the law of mass action, 58 reactions informed by Michaelis-Menten kinetics, 6 reversible reactions informed by Michaelis-Menten kinetics, 9 bi, 3 ping pong bi-bi kinetic reactions, and 31 rate laws that were either part of the old model or developed during its adaptation. Each functional form is underpinned by the experimental literature, and examples of all the functional forms are provided within the ordinary differential equations (**Appendix B**). In order to parameterise the kinetic reactions summarised in **Table A.3**, firstly a comprehensive search of the enzyme functional data repository, BRENDA was conducted (<http://www.brenda-enzymes.org/>) (Chang et al., 2015). Many of the enzymes identified within BRENDA had a wide range of values. Suitable values were selected based on considering the biological source of the kinetic information. Some of the model parameters could not be located within BRENDA, therefore to obtain these parameters, literature searches were conducted using PubMed, Science Direct and Google Scholar. The source of each parameter is indicated in column 4 of **Table A.4**. For a number of parameters, no kinetic information could be obtained, therefore several assumptions were used, the details of these are also given in **Table A.4**. The model was tuned until an 'optimal' set of parameters was obtained, which reflected the steady state values of key model outputs

Table 1. Data used to calculate initial values of cholesterol in each compartment. See table A.2 for abbreviations.

Species	Concentration	Molecular mass	Location Mass (kg)	Calculated mass (mg)	Reference
HFC	67.1nmol/mg	386.66	1.5	38,917.329	Mc Auley et al. (2012) & van de Pas et al. (2012)
HCE	18.5nmol/mg	649.08		18,011.97	
PFC	399.9mg/kg	-	64.8	25,913.52	
PCE	65.1mg/kg	-		4,218.48	

Utilising a plasma volume of 2.79L (van de Pas et al., 2012), the initial steady state values of VLDL-C, LDL-C and HDL-C (Sharma et al., 2010), could be converted to mg/plasma (**Table 2**). Lipoprotein values were converted back to mg/dL for analysis. When the model was simulated, VLDL-C, LDL-C and mean HDL-C values were consistent with these literature values (**Figure 4D**). Mean HDL-C was calculated from HDL₂ and HDL₃ and ndHDL-C subfractions. Cholesterol and unconjugated bile acids were steadily excreted at a rate of 0.1 and 0.05mg/hour respectively (**Figure 4E**). The initial steady state values for bile acids were assumed. For bile acids contained in the hepatic tissue, a value of 220mg was utilised for those conjugated (HCBA). A value of 180mg was assigned to represent the unconjugated proportion (HUBA), and a value of 200mg was chosen to represent HUBA-CoA. Within the ileocytes, a value of 50mg was utilised for both the conjugated and unconjugated forms (ICBA and IUBA). Within the intestinal lumen a steady state value of 200mg was assumed to represent conjugated bile acids, while a value of 50mg represented the steady state level of unconjugated bile acids (LCBA and LUBA). Levels remained constant over time (**Figure 4F**).

Table 2. Data used to calculate the initial values of lipoproteins. See Table A.2 for abbreviations.

Lipoprotein	Concentration (mg/dL)	Plasma Volume (L)	Calculated mass (mg)	Reference
VLDL-C	34	2.79	948.6	Sharma et al. (2010) & van de Pas et al. (2012)
LDL-C	115		3,208.5	
HDL-C	44		1,227.6	

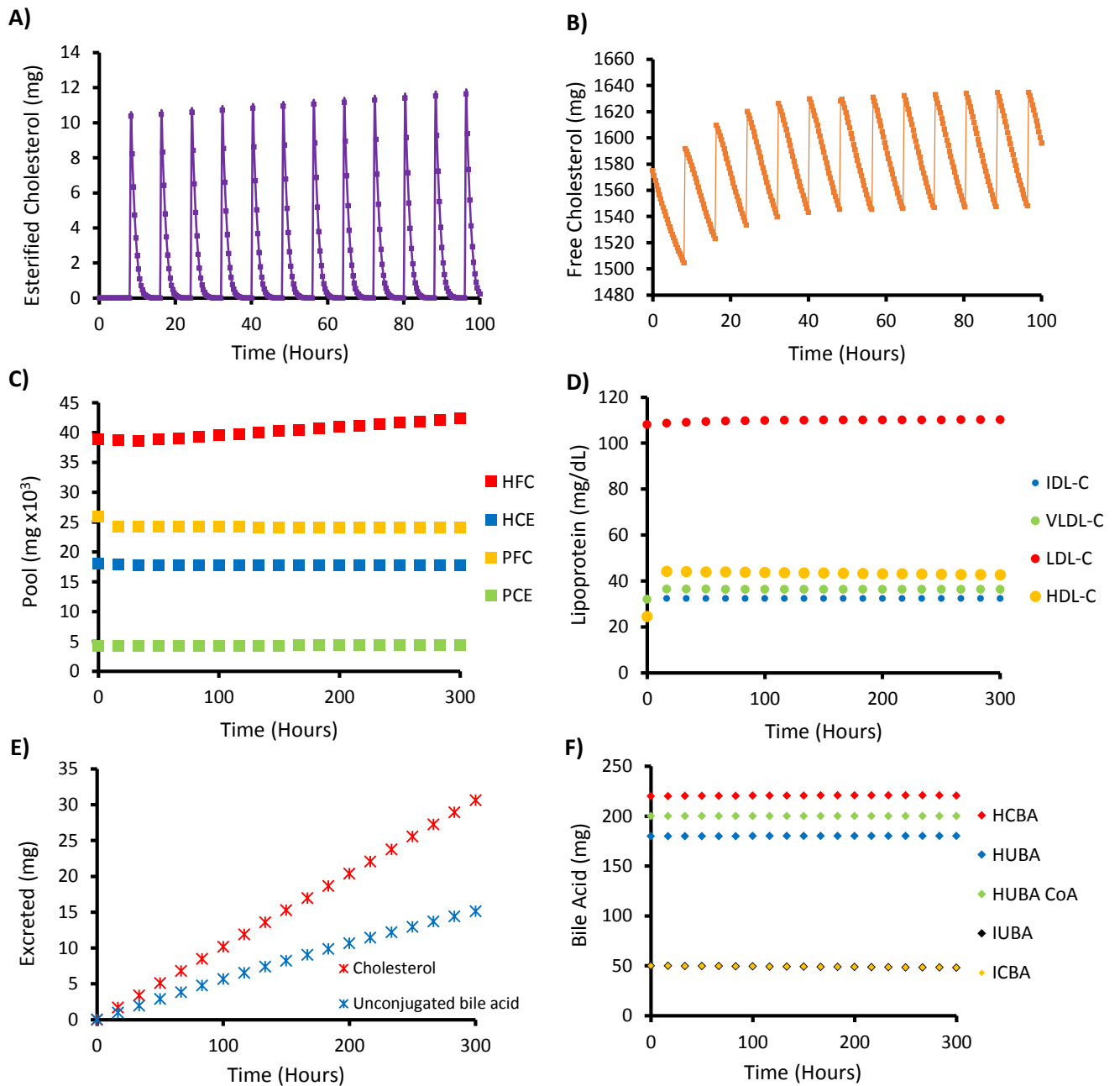


Figure 4. Model Outputs of **A)** ingestion of esterified cholesterol, **B)** jejuncyte pool of free cholesterol **C)** hepatic and peripheral pools of free and esterified cholesterol, **D)** lipoprotein profile **E)** excreted cholesterol and unconjugated bile acids and **F)** hepatic, ileocyte and intestinal lumen pools of conjugated and unconjugated bile acids. For D) mean HDL-C calculated from HDL₂, HDL₃ and ndHDL-C.

5.0. Results

5.1. Initial Examination of the Model

Initial simulations were performed to compare model output with our previous model. **Figure 5** illustrates how several key variables compare to the original model. Over a 15 day period it was found that the average concentration of LDL-C, mean HDL-C and total cholesterol reduced by 11%, 20% and 0.6% respectively, when compared to the original model. Mean HDL-C was calculated from HDL₂, HDL₃ and ndHDL-C. In contrast, IDL-C and VLDL-C increased by 33%

and 67% respectively in the updated model. The LDL-C:HDL-C ratio increased slightly from 2.52:1 in the original model to 2.77:1 in the updated model. Both of these ratios however are in line with current recommendations. A ratio <3.0:1 and <2.5:1 is optimal for males and females respectively, to reduce the risk of CVD onset (Millan et al., 2009).

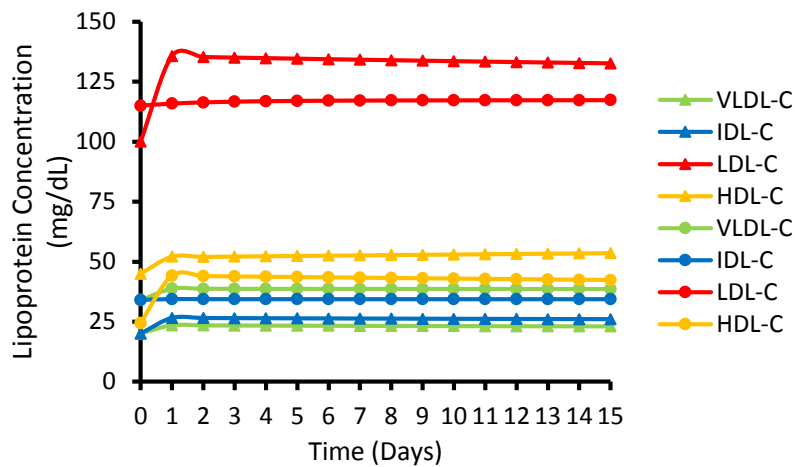


Figure 5. Comparison of the lipoprotein profile from the original and updated model. Model run for 15 days or equivalent (21,600 minutes). Circle markers represent the model output from the updated model. Triangle markers represent the lipoprotein profile from the Mc Auley et al. (2012) model. HDL-C from the updated model is the mean average of HDL₂, HDL₃ and ndHDL-C.

5.2. Dietary Cholesterol Ingestion

Individual differences exist in cholesterol metabolism from one individual to the next (Herron et al., 2003). This is emphasised by DC feeding studies which demonstrate notable differences in the response to DC from one individual to individual. Herron et al. (2003) demonstrated that hypo-responders could ingest ~640mg/day of cholesterol with no effect on LDL-C or HDL-C levels. In contrast, hyper-responders exhibited a 30% increase in LDL-C and an 8% increase in HDL-C (Herron et al., 2003). To determine if the model behaves as a hypo- or hyper-responder, fold increases of 0.5, 1, 2, and 3 were applied to both ingested free and esterified DC. To do this the events for DC ingestion were modified. For the 0.5-fold analysis, 44.335mg of free cholesterol and 6.3335mg of esterified cholesterol were ingested 3 times a day. Whereas an increase to 266.001mg of free cholesterol and 38.001mg of esterified cholesterol was ingested 3 times a day for a 3-fold dietary increase. A 2-fold increase in DC resulted in an increase of 0.0012% (0.0014mg/dL) in LDL-C, (>1000 hours). No change in mean HDL-C levels were observed (data not shown). A 2-fold increase in DC resulted in a 23.7% increase in HFC, and a negligible increase of 0.0002% in HCE. No difference in PFC or PCE was observed (**Figure 6A**).

5.3. Comparison with Clinical data

Both the original and updated models were compared to clinical data from a DC feeding study (**Figure 6B**). After a comprehensive literature search in this area, the most suitable time course data that could be obtained was from a healthy 31 year old normocholesterolaemic male that underwent an 11 week high cholesterol diet (1000mg/day) (Lin and Connor, 1980). The purpose of using this data, which clearly shows a hyper-response to DC, was to provide a comparison between this and our models. To compare the model to the clinical data, the intake of cholesterol was raised to 1000mg per day, and baseline LDL-C was reduced to 66.32mg/dL. To recreate a 1000mg/day cholesterol intake, the events were altered to reflect an intake of 291.667 FC and 41.667mg CE 3 times a day. Lin and Connor (1980) did not report LDL-C or HDL-C values, therefore LDL-C was calculated from total cholesterol and triglyceride values using the equation below, in which LDL-C can be calculated in the absence of HDL-C data (Anandaraja et al., 2005).

$$\text{Calculated LDL-C} = (0.9 \times \text{Total Cholesterol}) - \frac{0.9 \times \text{triglycerides}}{5} - 28$$

When the daily rate of cholesterol ingestion was raised, LDL-C increased from an initial value of 66.32mg/dL, to a steady state value of ~115mg/dL after one week. It is important to note this is the behaviour of LDL-C regardless of the rate of dietary cholesterol ingestion. This finding contrasts with the original model's response during this feeding regime, where LDL-C continues to rise after week one. It is feasible the behaviour of the updated model is biologically plausible, as several decades of experimental work in this area have suggested that cholesterol metabolism is tightly regulated in the majority of individuals, with hypo-responders known to exhibit a negligible lipoprotein response (Herron et al., 2003; Jones et al., 1996; Quintao et al., 1971). Consequently, it was important to establish which regulatory mechanisms responded effectively in the model to inhibit a rise in lipoprotein cholesterol. After examining the simulation data it was found a key regulatory mechanism which adapted to this increase in DC was cholesterol excretion (EFC). EFC increased by 25.1% by the end of the 11 week period. In addition to the compensatory effect of excreted cholesterol, intestinal and hepatic cholesterol also increased. It was found high DC resulted in the pool of CE in the intestinal lumen (LCE) increasing to 5.97mg after 11 weeks. This contrasted with a value of 0.15mg after a 'normal' cholesterol feeding regime over the same time period. LFC also increased by 4.75% as a result of this simulation. The amount of absorbed cholesterol, in the jejuncytes also increased; JFC and JCE levels were 24.41 and 0.03% higher respectively after the cholesterol feeding. In addition, HFC increased by 85.25%, while the increase in HCE was negligible (0.0005%). Bile acid levels and LDLr activity/number were unaffected during cholesterol feeding, which could account for the substantial cholesterol accumulation within the hepatic tissue, and marginal LDL-C response.

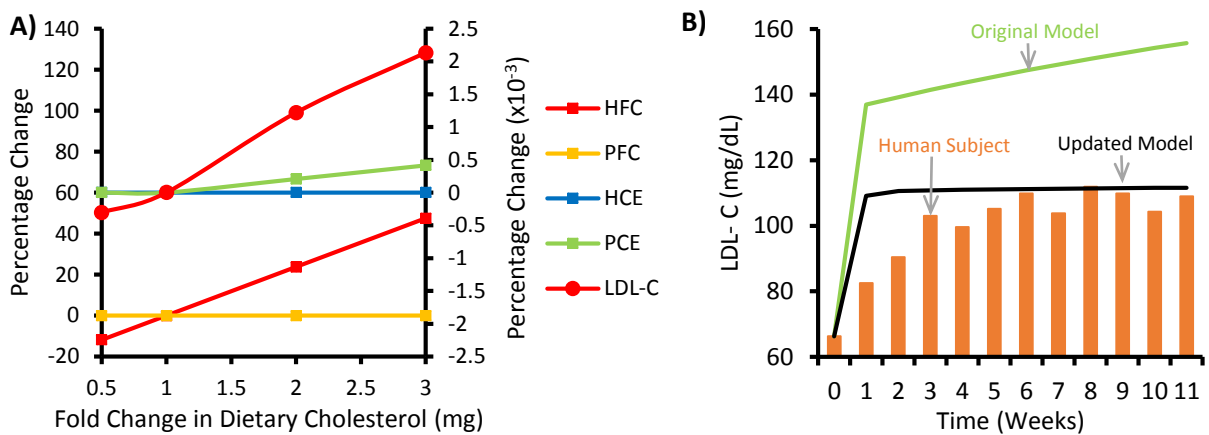


Figure 6. Cholesterol Ingestion Analysis. **A)** Response of LDL-C and cholesterol pools in the updated model to fold-change increases in dietary cholesterol after 1000 hours. HFC, hepatic free cholesterol; PFC, peripheral free cholesterol (left axis); HCE, hepatic cholesterol esters; PCE, peripheral cholesterol esters; and LDL-C (right axis). **B)** Goodness of fit of model to clinical data. Comparison of LDL-C concentration in response to simulated cholesterol feeding in the Mc Auley et al. (2012), and the updated model, with a human cholesterol feeding study. A 1000mg dose of cholesterol was ingested per day by a human subject and induced in both models for an 11 week trial period and the LDL-C monitored. Baseline LDL-C of both models was set at 66.32mg/dL to match clinical data. Black line represents updated model the green line represents the Mc Auley et al. (2012) model, and the orange bar represents the LDL-C response to cholesterol feeding in a human subject (Lin and Connor, 1980).

5.4. Acute daily ingestion of saturated fat

Diets high in SFA are associated with raised plasma cholesterol levels. High levels of SFA lead to an increase in LDL-C by three possible mechanisms. 1) LDL-C clearance rate, as SFA

suppresses LDLr activity (Woollett et al., 1992). 2) There is an increase in the rate of cholesterol synthesis (Jones et al., 1994). 3) Modulation of RCT occurs (Berard et al., 2004; Jansen et al., 2000). RCT is affected in 2 ways. Firstly, CETP is upregulated, thus transporting esterified cholesterol from HDL-C to LDL-C and VLDL-C at an increased rate (Jansen et al., 2000). Secondly, LCAT is inhibited which reduces the esterification of cholesterol in HDL-C. This reduces the concentration gradient for the transfer of cholesterol from the peripheral tissues to HDL. To replicate SFA intake variations, 7 parameters were altered simultaneously (**Table 3**). This analysis resulted in increases to LDL-C, HCE and PCE levels, whereas HDL-C decreased (**Figure 7**). LDL-C, HCE, and PCE increased 1.7, 0.4 and 0.5% respectively for the 2 fold-analysis. There was no effect on hepatic and peripheral free cholesterol (HFC and PFC) after 1000 hours.

Table 3. Summary of parameters involved in fold analysis of SFA intake.

Reaction	Parameter	Fold Change				
		0.5	1	2	3	
36	LCAT activity	K_{hdl3}	8×10^{-4}	4×10^{-4}	2×10^{-4}	1.33×10^{-4}
39	CETP mediated transfer to VLDL	K_{cetp2}	5×10^{-8}	1×10^{-7}	2×10^{-7}	3×10^{-7}
40	CETP mediated transfer to LDL	K_{cetp1}	5×10^{-8}	1×10^{-7}	2×10^{-7}	3×10^{-7}
60	LDLr degradation	K_1	5×10^{-8}	1×10^{-7}	2×10^{-7}	3×10^{-7}
78	Hepatic acetyl CoA synthesis	K_1	0.5	1	2	3
96	Peripheral acetyl CoA synthesis	K_1	5×10^{-6}	1×10^{-5}	2×10^{-5}	3×10^{-5}
114	Intestinal acetyl CoA synthesis	K_1	0.5	1	2	3

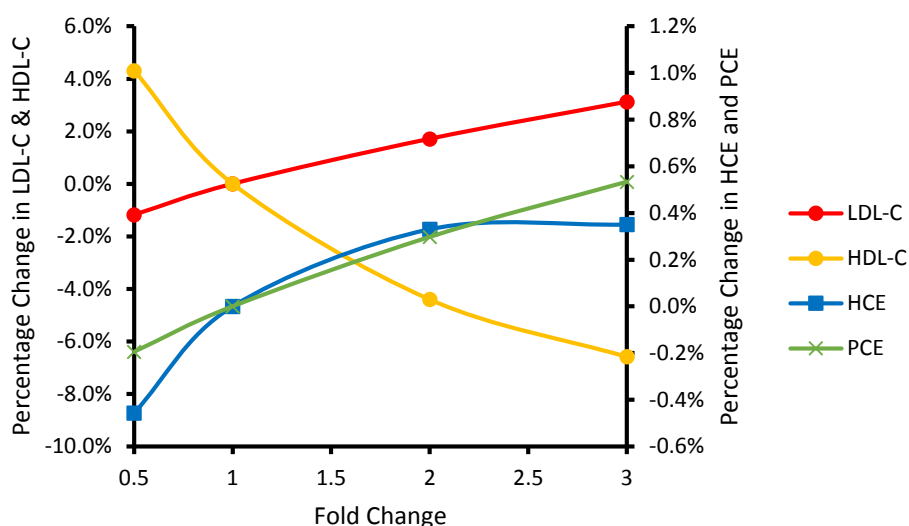


Figure 7. Fold change analysis of dietary SFA. The percentage change in LDL-C, HDL-C (left axis), HCE and PCE (right axis) monitored with fold changes to SFA intake. A total of 7 parameters were modified in the fold change analysis to represent an equivalent change in dietary SFA. To represent an increase in dietary SFA, increases to the rate of LDLr degradation, cholesterol synthesis and CETP mediated transport, and decreases to the LCAT mediated cholesterol esterification were conducted. Measurements recorded at 1000 hours.

5.5. Local Sensitivity Analysis

To gain further insight into the robustness of the model a sensitivity analysis (SA) was performed. A robust model is resilient to relatively small perturbations in parameter values. Therefore given the lack of sensitivity of our model during our initial investigations the aim of the SA was to investigate the effect of both local and global parameter changes on model outputs, in particular LDL-C, and to identify critical parameter inputs. We firstly completed a local SA and calculated the sensitivity coefficients (S_i) for each of the parameters of the model. To do this we removed the discrete feeding events and replaced them with a continuous feed for both DC and DFC. **Appendix C** gives the percentage change in output values for a 1% change in each of the 266 model parameters and ranks them by sensitivity, while **Figure 8** presents the 10 most sensitive parameters of this analysis.

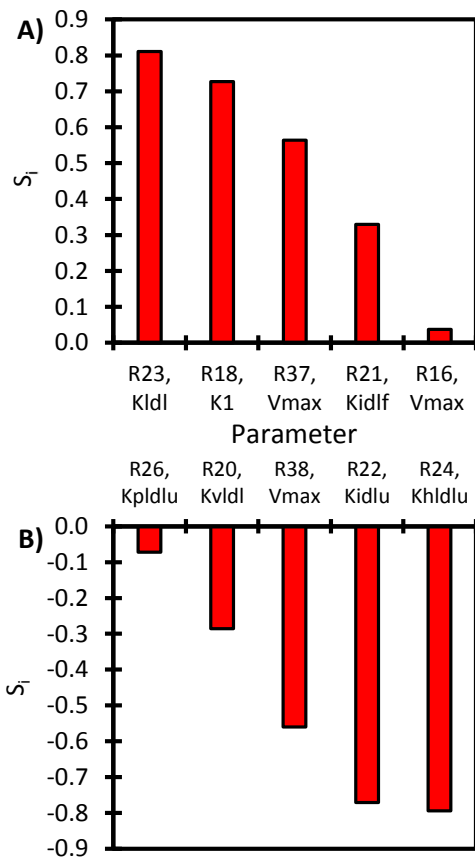


Figure 8. The 10 most sensitive parameters. S_i , sensitivity index, percent change in LDL-C for a 1% change in model parameters; R, reaction. See Table A.3 for reaction details.

5.6. Global analysis

Given the uncertainty surrounding a number of the parameter values, especially those associated with the less well characterised aspects of cholesterol metabolism, it is apparent that any inferences from our local sensitivity analysis should be interpreted with caution. Therefore, we deemed that it would be appropriate to conduct a global sensitivity analysis. A global sensitivity analysis attempts to mitigate this problem by examining model parameters in broader regions of parameter space. The rate constant for hepatic LDLr degradation was selected as our parameter of interest for the global scan (reaction 60 in Table A.3). This parameter was selected as the experimental literature suggests this is an important parameter for maintaining cholesterol homeostasis, and the reduction in hepatic LDLr numbers and/or activity has been associated with intrinsic ageing (Field and Gibbons, 2000). Furthermore, of the mechanisms associated with ageing, this parameter was most sensitive (see appendix C). As no *priori* data was available on this parameter, it was determined that an appropriate sampling choice would be based on a uniform probability density function (Marino et al., 2008). **Figure 9** presents the results from a global sensitivity analysis, where random samples were taken over a range of a 2 fold increase and decrease in the default value for this rate constant, with A) all “known” parameters scanned and unknown parameters fixed, B) all unknown parameters scanned and “known” parameters fixed, and C) all parameters scanned simultaneously, at 250 hours (N=500). See appendix C for a list of the scanned parameters for each of these simulations. Each scanned parameter had random samples taken between a 2 fold increase and decrease of its default value. The variability of LDL-C was small when 151 “known” parameters were scanned, and 113 unknown parameters remained fixed. The results show that LDL-C varied ~2mg/dL for any value of the rate constant for hepatic LDLr degradation between a 2 fold increase and decrease from its default value (**Figure 9A**). In contrast, LDL-C varied by ~500mg when the “known” parameters were fixed and the unknown parameters scanned. Consequently the relationship between hepatic LDLr degradation and LDL-C was lost (**Figure 9B**). When all 266 parameters were scanned (including the 2 parameters used for the counter reactions for DFC and DCE intake), the range of LDL-C values remained similar. A marginal further reduction in R^2 was also observed (**Figure 9C**). The significant difference in LDL-C variation when scanning parameters with well characterised kinetics versus parameters with assumed parameter values suggests the unknown parameters influence the behaviour of the model significantly. Moreover, the large range in LDL-C when unknown parameters were scanned, reinforces our finding from the local sensitivity scan, that parameters with assumed kinetics were most sensitive.

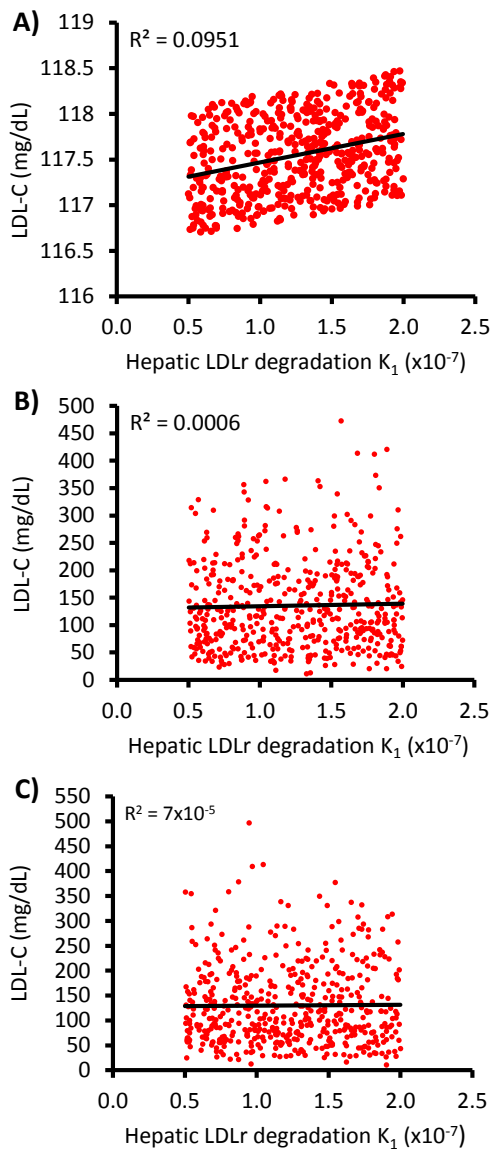


Figure 9. Global sensitivity analysis of parameter K_1 , hepatic LDLr degradation. Role of parameter K_1 for hepatic LDLr degradation (reaction 60) with A) all unknown, assumed, and adapted parameters fixed, B) all “known” parameters fixed, and C) all parameters scanned. Non-fixed parameters were analysed between a 2-fold increase and decrease of that parameter value; calculated at 250 hours, $N=500$. R^2 = Coefficient of determination. See section 5.6 for details of each figure.

5.7. Ageing

As previously discussed, the ageing process affects many of the key mechanisms involved in cholesterol metabolism. Specifically, there is 1) an age-related decline in hepatic LDLr and subsequent reduction in LDL-C clearance (Millar et al., 1995), 2) an increase in cholesterol absorption (Duan et al., 2006), 3) a decline of BSH-positive intestinal microflora, such *Lactobacillus* and *Bifidobacterium* species (Hopkins and Macfarlane, 2002), and 4) a decline in bile acid synthesis (Bertolotti et al., 2007). To represent the ageing process, 4 parameters were modified. These modifications were a 2-fold decrease to the rate of NPC1L1 degradation, bacterial deconjugation, and CYP7AI activity. In addition, the rate of LDLr degradation was increased 2-fold (**Table 4**). The inverse of this analysis, was assumed to represent a young individual.

Table 4. Summary of parameters involved in the ageing analysis.

Reaction	Parameter	Age group		
		Young	Default	Old
45 Bile acid Deconjugation	V_{max}	1×10^{-5}	5×10^{-6}	2.5×10^{-6}
54 NPC1L1 degradation	K_1	2×10^{-7}	1×10^{-7}	5×10^{-8}
60 LDLr degradation	K_1	5×10^{-8}	1×10^{-7}	2×10^{-7}
132 CYP7AI activity	V_{max}	1.11358×10^{-3}	5.5679×10^{-4}	2.78395×10^{-4}

By simultaneously conducting a 2-fold decrease to the parameters K_{cept1} and K_{cept2} , a CETP genotype such as I405V could also be replicated (reactions 40 and 39 respectively, Table A.3). A 2 fold increase in K_{cept1} and K_{cept2} was also conducted to replicate a genotype reflective of a genetic predisposition for atherogenesis. These CETP genotypes were applied to each age category and LDL-C was recorded after 1000 hours (**Figure 10**). The ageing process resulted in an increased LDL-C for all CETP genotypes. The ageing process, in the presence of the genotype representing high CETP activity, resulted in a 1.6% increase in LDL-C. Similarly, the normal CETP genotype also produced a 1.6% increase in LDL-C with ageing. The CETP I405V genotype however resulted in a less significant rise (0.6%) in LDL-C with the ageing process. By contrasting the genotypes within each age category, the beneficial effects of the CETP I405V genotype were further substantiated. This advantageous genotype was more obvious in the aged category than the youth and normal categories. A 1.63% reduction in LDL-C was observed in the aged category, compared to a 0.64% reduction in both the normal and youth category. On the contrary, the effect of a genetic predisposition to atherosclerosis, was more apparent in the youth and normal categories, than the aged category, although this was negligible (0.617, 0.617 and 0.616% respectively).

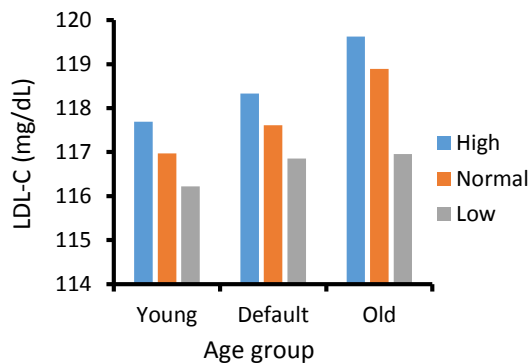


Figure 10. Role of CETP genotypes on LDL-C with ageing. To mimic an aged individual, a 2 fold increase in LDLr degradation, and a 2 fold decrease in NPC1L1 degradation, bile acid deconjugation, and CYP7A1 activity was applied. For the youth analysis, the inverse of this protocol was conducted. A 2 fold increase or decrease in CETP activity was then applied to each group. High (blue), normal (orange), and low (grey) CEPT activities examined. LDL-C measured at 1000 hours.

6.0. Discussion

We have updated a whole-body mathematical model of cholesterol metabolism. The updated model includes 96 additional mechanisms, and has been deposited in the BioModels database (MODEL1508170000). In contrast to the original model, this model is significantly less sensitive to cholesterol feeding, and can be categorised as a hypo- rather than hyper-responder to cholesterol ingestion. This finding is perhaps unsurprising in light of experimental evidence which suggests 62.5% of healthy normocholesterolaemic males exhibit a hypo-response to dietary cholesterol (Herron et al., 2003). Moreover, it is widely regarded that cholesterol metabolism adapts to cholesterol feeding by invoking two major compensatory regulatory mechanisms, these are an increase in cholesterol excretion, and a decrease in whole-body cholesterol biosynthesis (Quintao et al., 1971). When there is effective feedback control mechanisms this can result in dietary cholesterol failing to illicit a response in lipoprotein metabolism. For instance, McNamara et al. (1987) demonstrated using 12 week studies that both low and high levels of dietary cholesterol did not induce significant increases in mean levels of plasma total, LDL, or HDL cholesterol. Moreover, there was no relationship between baseline plasma cholesterol levels and sensitivity to a dietary cholesterol challenge (McNamara et al., 1987).

In our model, clearly LDL-C did not rise, however both hepatic free cholesterol and intestinal cholesterol increased significantly during high cholesterol feeding. In the seminal work by Quintao et al. (1971) it is postulated that cholesterol feeding in humans is primarily controlled by one compensatory mechanism – an increase in cholesterol excretion, but not bile acid excretion, our model helps to substantiate this claim. Intriguingly, the authors also suggest that when there is an accumulation of cholesterol in the body pools, the effects on lipoprotein levels are insignificant, regardless of the quantity of cholesterol absorbed. This appears to be the case in humans, as regulatory processes inhibit an increase in lipoprotein cholesterol when body pools accumulate. Model organisms have also shown that body pools of cholesterol can increase significant as a result of cholesterol feeding. Ohtani et al. (1990) suggest that, in hamsters, an additional 0.1% dietary cholesterol is enough to induce the accumulation of cholesterol in the liver through the chylomicron remnant pathway (Ohtani et al., 1990). Wang et al. (2010) also reported that concentrations of the hepatic total cholesterol increased ~4-fold by with a 1% increase in dietary cholesterol in rats (Wang et al., 2010).

Although the increases in the pools of cholesterol in our model, is similar to what has been reported experimentally, this could also suggest that our regulatory mechanisms may not be sufficiently adequate to deal with the excess cholesterol. One mechanism in this model that may need to be improved is the turnover of LDLr. In the updated model, an increase in dietary cholesterol, did not elicit a decrease in LDLr number. This contrasted with the previous model. This is an important point that needs to be emphasised, Hennessey et al. (1992) have demonstrated that a high intake of dietary cholesterol suppresses hepatic LDLr mRNA by 29% in cebus monkeys (Hennessey et al., 1992). Conversely, Spady and Cuthbert (1992) observed that rats on fed high levels of dietary cholesterol, did not exhibit suppressed hepatic LDLr activity despite a 350-fold increase in hepatic cholesterol ester levels (Spady and Cuthbert, 1992). More recently, Boone et al. (2011) found that LDLr expression in Sprague-Dawley rats was not altered by dietary cholesterol. There was also a significant increase in mean hepatic cholesterol and paradoxically a decrease in serum cholesterol (Boone et al., 2011). However, other studies in humans, such as Lichtenstein et al. (1993) demonstrate that the addition of dietary cholesterol (~1.5 eggs per/day) to a corn oil-diet, can result in a significant increase in concentrations of total cholesterol (6%), LDL-C (8%), and HDL-C (7%), when compared to diets with low dietary cholesterol, in 14 middle-aged and older individuals. Studies such as this consolidate the view that LDLr activity is downregulated in response to cholesterol feeding. Despite the lack of sensitivity of lipoprotein metabolism to cholesterol feeding, our model was sensitive to increases in dietary SFA. This finding is in line with the experimental literature as dietary SFA has been shown clinically to have a more profound effect on cholesterol metabolism than dietary cholesterol (He and Fernandez, 1998; Herron et al., 2003).

Large inter-individual differences in cholesterol metabolism, can in part be explained by genetic factors. For example, the I405V genotype is associated with longevity and reduced levels of CETP. This results in larger LDL-C and HDL-C particle size, in addition to reduced levels of LDL-C and increased HDL-C. This genotype is related to lower prevalence of hypertension, CVD, diabetes mellitus and metabolic syndrome (Atzmon et al., 2005; Barzilai et al., 2003). The role of a CETP genotype for reduced CETP, such as I405V, in reducing the risk of age-associated disease, was consolidated with our simulations. It was demonstrated that increased CETP activity resulted in raised LDL-C in all age groups. Conversely, decreased CETP activity, led to a lower LDL-C in all age categories. Furthermore, the age-related increase in LDL-C was reduced with the presence of the I405V genotype (0.6% vs. 1.6%). Although it is important to note that a reduction in CETP levels and subsequent increase in HDL-C, either due to a genetic polymorphism or pharmaceutical intervention, has been associated with an increase in mortality rate (Barter et al., 2007; Zhong et al., 1996). It is therefore important that the findings from this work, and other studies, are fully evaluated before *in vivo* application.

An important finding from our updated model was that the parameter values associated with the less well characterised aspects of cholesterol metabolism were particularly sensitive to changes in their values. Moreover, the cholesterol biosynthesis pathway, which has been well characterised, was a particularly robust part of the model. These findings from our local sensitivity analysis were supported by our global sensitive analysis, which demonstrated the increase in variation in LDL-C when unknown parameters were scanned, compared with known parameters. Therefore, we also have to consider the fact that the functional forms we have used to represent the interactions in our model may need further improvements, and that interactions may be due to many components of the model, such as RCT, lipoprotein dynamics and the enterohepatic circulation of bile acids, being poorly understood from a kinetic perspective.

There are a number of aspects of the model that could be developed in the future. For example, when conjugated and unconjugated bile acid levels are in a steady state, the V_{max} value for the deconjugation of bile acids was significantly lower than that reported in the experimental literature (reaction 45 Table A.3). For example, for the intestinal bacteria *Clostridium perfringens*, *Bacteriodes fragilis* and *Lactobacilli* have suggested V_{max} values of 0.05, 38.4 and 25.4mg/min/mg. However, a far lower V_{max} value of 5×10^{-6} was used in the model (Gopal-Srivastava and Hylemon, 1988; Lundeen and Savage, 1992; Stellwag and Hylemon, 1976). Areas such as cholesterol and bile acid synthesis could also be further developed to incorporate less well understood mechanisms that have not been included in the model. Furthermore, this model could be combined with other pre-existing models, or further developed to include other

compartments such as the gallbladder, or other interconnected metabolic pathways such as fatty acid or carbohydrate metabolism, or cortisol homeostasis. For example, cortisol is synthesised from cholesterol and is involved in provoking the breakdown of lipids, and a wide variety of other metabolites, therefore this model could be connected to a previously developed computational model of whole-body cortisol metabolism (Mc Auley et al., 2013; McAuley et al., 2009).

7.0. Conclusions

In combination with our previous model, the updated model presented here emphasises the important role of intrinsic ageing in disrupting whole-body cholesterol metabolism. Constructing this model has revealed a number of findings. Firstly, simulations indicated that the model is a hypo-responder to cholesterol feeding, but is sensitive to simulated SFA feeding. Secondly, the model tentatively supports that genotypes such as I405V has a protective role in healthy ageing. The model also highlights the robustness of the cholesterol biosynthesis pathway, and suggests key areas where experimental work needs to focus in the future. This study has highlighted the need for kinetic data in several key areas of cholesterol metabolism, including RCT, lipoprotein dynamics and the enterohepatic circulation of bile acids. As these areas could provide potential therapeutic targets for lowering cholesterol, we believe that a more in depth understanding of this process could lead to novel interventions in the future. Finally, this work has demonstrated the important role computational systems biology has to play in improving our understanding of lipid metabolism and health-span.

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