Comparative Hepatic and Intestinal Metabolism and Pharmacodynamics of Statins

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ABSTRACT

This study aimed to comprehensively investigate the in vitro metabolism of statins. The metabolism of clinically relevant concentrations of atorvastatin, fluvastatin, pitavastatin, pravastatin, rosuvastatin, simvastatin, and their metabolites were investigated using human liver microsomes (HLMs), human intestine microsomes (HIMs), liver cytosol, and recombinant cytochrome P450 enzymes. We also determined the inhibitory effects of statin acids on their pharmacological target, 3-hydroxy-3-methylglutaryl–coenzyme A (HMG-CoA) reductase. In HLMs, statin lactones were metabolized to a much higher extent than their acid forms. Atorvastatin lactone and simvastatin (lactone) showed extensive metabolism [intrinsinc clearance (Clint) values of 3700 and 7400 μl/min per milligram], whereas the metabolism of the lactones of 2-hydroxyatorvastatin, 4-hydroxyatorvastatin, and pitavastatin was slower (Clint 20–840 μl/min per milligram). The acids had Clint values in the range <0.1–80 μl/min per milligram. In HIMs, only atorvastatin lactone and simvastatin (lactone) exhibited notable metabolism, with Clint values corresponding to 20% of those observed in HLMs. CYP3A4 and 2C9 were the main statin-metabolizing enzymes. The majority of the acids inhibited HMG-CoA reductase, with 50% inhibitory concentrations of 4–20 nM. The present comparison of the metabolism and pharmacodynamics of the various statins using identical methods provides a strong basis for further application, e.g., comparative systems pharmacology modeling.

SIGNIFICANCE STATEMENT

The present comparison of the in vitro metabolic and pharmacodynamic properties of atorvastatin, fluvastatin, pitavastatin, pravastatin, rosuvastatin, and simvastatin and their metabolites using unified methodology provides a strong basis for further application. Together with in vitro drug transporter and clinical data, the present findings are applicable for use in comparative systems pharmacology modeling to predict the pharmacokinetics and pharmacological effects of statins at different dosages.

Introduction

3-Hydroxy-3-methylglutaryl–coenzyme A (HMG-CoA) reductase inhibitors (statins) are widely used in the treatment of hypercholesterolemia. Although statin drugs are effective and safe in most patients, many users experience poor efficacy or adverse drug reactions (Pazzucconi et al., 1995; Yebyo et al., 2019). The muscle toxicity of statins is a dose- and concentration-dependent phenomenon (Bradford et al., 1991; Dujovne et al., 1991), and the risk of toxicity increases along with the plasma concentrations of statins. This may be caused by drug-drug interactions or inherited defects in proteins affecting statin disposition, such as CYP2C9 and 3A4, organic anion transporting polypeptide (OATP) 1B1, or breast cancer resistance protein (Thompson et al., 2003; Graham et al., 2004; Neuvonen et al., 2006; Pasanen et al., 2006; Keskiörala et al., 2009b). Both acid and lactone forms of statins can interact with proteins involved in drug disposition. Lovastatin and simvastatin are administered as lactone prodrugs, whereas other statins are given in the active acid form. In the body, however, significant amounts of many statins are converted to their corresponding acid/lactone form (Neuvonen et al., 2006). Cytochrome P450 enzymes metabolize the lipophilic statin lactones more rapidly than statin acids (Fujino et al., 2004). Indirectly, uridine diphosphate-glucuronosyltransferase (UGT) enzymes may participate in lactone formation by mediating glucuronidation of statin acids. The acyl glucuronides formed in this process are converted to their corresponding acid/lactone form (Neuvonen et al., 2006). This work was supported by the European Research Council Consolidator Grant [Grant Agreement 725249]. The authors declare no conflict of interest.

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ABBREVIATIONS: Clint, intrinsic clearance; fub, unbound fraction in microsomes; HLM, human liver microsome; HLC, human liver cytosol; HIM, human intestine microsome; HMG-CoA, 3-hydroxy-3-methylglutaryl–coenzyme A; LC-MS/MS, liquid chromatography-tandem mass spectrometry; OATP, organic anion transporting polypeptide; PAPS, 3'-phosphoadenosine 5'-phosphosulfate; PBPK, physiologically based pharmacokinetic; PMSF, phenylmethylsulfonyl fluoride; SULT, sulfotransferase; UDPGA, uridine 5'-diphospho-glucuronic acid; UGT, uridine diphosphate-glucuronosyltransferase.
CYP2C9 (Fischer et al., 1999; Hirvensalo et al., 2019). Pitavastatin, pravastatin, and rosuvastatin are excreted mainly unchanged (Neuvonen et al., 2006). Hence, the available statins differ significantly in their pharmacokinetic characteristics and susceptibility to altered metabolizing enzyme function.

Interestingly, the pharmacological target of statins, HMG-CoA reductase, is expressed in the same location as the cytochrome P450 enzymes, the endoplasmic reticulum of hepatocytes (Corsini et al., 1995). Thus, this enzyme is also present in human liver microsomes (HLMs). Similar to cytochromes P450, the active site of HMG-CoA reductase faces the cytoplasm, and the enzyme uses NADPH for its catalytic activity (Corsini et al., 1995). Statin acids, which are structurally similar to its substrate HMG-CoA, competitively inhibit it in the nanomolar range, leading to a reduction of the HMG-CoA-mevalonate pathway. Also, several statin metabolites have been depicted to be inhibitors of this reaction, but in vitro data for, e.g., the 2- and 4-hydroxy metabolites of atorvastatin do not seem to be publicly available.

Many published physiologically based pharmacokinetic (PBPK) models of statins can be found in the literature, primarily aiming to evaluate their drug-drug interaction and pharmacogenetic properties. During the past two decades, PBPK modeling has evolved as an important tool in drug development, regulatory review, and clinical study design (Sager et al., 2015). PBPK modeling also holds the potential to become a valuable tool to inform drug and dosing selection in clinical practice (Johnson and Rostami-Hodjegan, 2011; Jamei, 2016; Venkatakrishnan and Rostami-Hodjegan, 2019). For such purposes, it is crucial that the data used for model development of a set of drugs are comparable and high in quality. Although the metabolism of statins has been widely investigated in vitro, only a few substrate depletion studies comparing the microsomal metabolism of different statins have been published (Fujino et al., 2004; Gertz et al., 2010, 2011; Varna et al., 2014). When appropriately applied, the substrate depletion approach results in intrinsic clearance (CL_{int}) values, which include all microsomal metabolic pathways of the tested compound (Obach, 1999), and can be scaled to hepatic metabolic clearance for PBPK modeling. Hence, to obtain comparable estimates of the metabolism of the widely used statins atorvastatin, fluvastatin, pitavastatin, pravastatin, rosuvastatin, and simvastatin and their metabolites, we carried out an extensive in vitro study in subcellular hepatic and intestinal fractions as well as in recombinant cytochrome P450 enzymes using low, clinically relevant statin concentrations. Moreover, to simulate the pharmacological response of statins in PBPK models, unbound statin concentrations in hepatocytes can be linked to in vivo measurements of their pharmacodynamic potency. Therefore, we also compared the inhibitory effects of these statins on their pharmacological target HMG-CoA reductase.

Materials and Methods

Chemicals and Reagents. Atorvastatin (acid), atorvastatin lactone, 2-hydroxyatorvastatin (acid), 2-hydroxyatorvastatin lactone, 4-hydroxyatorvastatin (acid), 4-hydroxyatorvastatin lactone, mevalonate lactone, pravastatin (acid), rosuvastatin (acid), and all internal standards (Supplemental Table 1) were purchased from Toronto Research Chemicals (Toronto, Canada). 3R,5S-fluvastatin (acid), 3S,5R-fluvastatin (acid), pitavastatin (acid), and pitavastatin lactone were obtained from Santa Cruz Biotechnology (Dallas, Texas), and simvastatin acid and simvastatin (lactone) from SynFine Research (Ontario, Canada). Adenosine 3-phosphate 5'-phosphosulfate triethylammonium salt (PAPS), alamethicin, NADPH, uridine 5'-diphospho-glucuronic acid (UDPGA), and HMG-CoA reductase assay kits were obtained from Sigma-Aldrich (St. Louis, MO). HLMs (XTreme 200, a pool of 200 mixed-sex donors), human liver cytosol (HLC: XTreme 200, a pool of 200 mixed-sex donors), and human intestine microsomes (HIMs), both normal (a pool of 15 mixed-sex donors) and free from phenylme-thylsulfonyl fluoride (PMSF; optimal in carboxylesterase activity; a pool of six mixed-sex donors), were purchased from Sekisui XenoTech (Tokai, Japan). The following recombinant EasyCYP Bactosomes were obtained from Cypex Ltd (Dundee, UK): CYP1A2R, CYP2A6BR, CYP2B6BR, CYP2C8BR, CYP2C9BR, CYP2C19BR, CYP2D6R, CYP2E1BR, CYP2J2LR, CYP3A4BR, and CYP3A5BLR. Other chemicals were from Merck (Darmstadt, Germany).

Metabolism in Human Liver Microsomal Incubations. The metabolic depletion of statins was first measured in HLMs. With the exception for buffer controls, all incubations contained substrate, microsomes (0.2 mg/ml) in sodium phosphate buffer (0.1 M, pH 7.4) with MgCl2 (5 mM). For simvastatin, the protein concentration was reduced to 0.1 mg/ml after the initial experiment. The depletion of each parent statin and its corresponding acid/lactone was studied in four different conditions: 1) addition of NADPH (1 mM) to the reaction mixture to measure the cytochrome P450–mediated metabolism; 2) addition of NADPH (1 mM) and UDPGA (5 mM) to measure both cytochrome P450– and UGT-mediated metabolism; 3) no addition of cofactors (negative control); and 4) no addition of cofactors or microsomes (buffer control). In condition 2, alamethicin (11 μg/ml) was also included in the incubations to allow pore formation. The metabolism of the 2- and 4-hydroxy metabolites of atorvastatin were studied using conditions 1, 3, and 4. The initial incubation concentrations of each substrate are listed in Supplemental Table 2 and Table 1.

All incubations were carried out once in triplicate on 96-well plates. The mixtures of substrate and microsomes (or only substrate in buffer in condition 4) were first preincubated for 15 minutes at 37°C and 350 rpm. After preincubation, cofactors were added to initiate the reactions in conditions 1 and 2. Samples were taken at 0, 5, 10, 20, 40, and 60 minutes. For atorvastatin lactone and simvastatin, the sampling times were shorter (0.5, 1, 3, 5, 7, and 10 minutes). Reactions were stopped by diluting samples 1:3 with ice-cold acetonitrile containing internal standard (1:2 for rosuvastatin) (Supplemental Table 1) and further handled as described below in the section Sample Processing and Analysis of Metabolism Samples.

Metabolism in Human Intestine Microsomal Incubations. The metabolic depletion of parent statins and their corresponding acid/lactone was also measured in HIMs. With the exception of buffer controls, all incubations contained substrate, microsomes (0.2 mg/ml) in sodium phosphate buffer (0.1 M, pH 7.4) with MgCl2 (5 mM). For each substrate (incubation concentrations listed in Supplemental Table 2 and Table 1), the depletion was studied in four different conditions: 1) addition of NADPH (1 mM) and UDPGA (5 mM) to measure both cytochrome P450– and UGT-mediated metabolism (alamethicin 11 μg/ml also included), 2) no addition of cofactors (negative control), and 3) no addition of cofactors nor microsomes (buffer control). Furthermore, to screen for the potential effects of intestinal carboxylesterases on the metabolism of statins, each statin was also incubated with 4) 0.2 mg/ml PMSF-free HIMs (optimal in esterase activity). No cofactors were added to PMSF-free HIM incubations.

All incubations were carried out once in triplicate, preincubated, and stopped in a similar manner as described for the HLM incubations above. In conditions 1–3, samples were taken at 0, 5, 10, 20, and 45 minutes. In condition 4, only two samples were taken: at 0 and 25 minutes.

Metabolism Screening in Human Cytosolic Incubations. To screen for statin metabolism mediated by cytosolic enzymes, each statin was incubated with HLC (0.2 mg/ml) in sodium phosphate buffer (0.1 M, pH 7.4) with MgCl2 (5 mM) at 37°C and 350 rpm, either 1) without cofactors or 2) with the sulfotransferase (SULT) cofactor PAPS (100 μM). Two samples were taken: 0 and 25 minutes. These conditions were selected to match those of the HIM screening (test setting 4 above). All incubations were carried out once in triplicate, and reactions were stopped as above.

Metabolism Screening with Recombinant Cytochrome P450 Enzymes. The depletion of statins was also measured in recombinant cytochrome P450 incubations. In the screening, the substrates were incubated with each one of 11 cytochrome P450 isoforms (CYP1A2, CYP2A6, CYP2B6, CYP2C8, CYP2C9, CYP2C19, CYP2D6, CYP2E1, CYP2J2, CYP3A4, and CYP3A5) separately at a protein concentration of 0.2 mg/ml. The substrate concentrations were the same ones as in the HLM depletion experiment (Supplemental Table 2; Table 1). Samples were collected at 0, 30, and 60 minutes. All incubations were carried out once in triplicate, and reactions were stopped as above.

Inhibition of Metabolism in Microsomal Incubations. To confirm the results from the recombinant cytochrome P450 screening, inhibition studies in HLMs and HIMs were carried out for selected statins (HLMs: atorvastatin and atorvastatin lactone, 3R,5S- and 3S,5R-fluvastatin, pitavastatin lactone, 2-hydroxy- and 4-hydroxy metabolites of atorvastatin were not publicly available.

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Calculated and measured in vivo plasma concentration data, and initial incubation concentrations (C₀) of the statins tested.

Fraction unbound in plasma and peak concentration data in plasma and blood-to-plasma concentration ratios were collected from the University of Washington Drug Interaction Database (September 21, 2020) or from ADMET Predictor (version 10; Simulations Plus, Lanchester, CA). The concentration data of the fluvastatin enantiomers are from Hirvensalo et al. (2019), and those of the atorvastatin metabolites from Kantola et al. (1998) (single-dose studies).

| Compound                | BP  | ℓ_{up} | C_{max} | C_{gut} | C_{max} | C_{ent} | C_{max, portal} | C_{max, portal} | C_{0} in HLMs, HLC, rCYPs | C₀ in HIMs |
|-------------------------|-----|--------|---------|---------|---------|---------|-----------------|-----------------|--------------------------|---------------|
| Atorvastatin            | 0.667 | 0.02 | 0.31 (40 mg) | 0.00062 | 290 | 8.7 | 0.69 | 2.5 | 0.051 | 0.05 | 1 |
| Atorvastatin lactone    | 0.716 | 0.04 | 0.0070 (40 mg) | 0.00028 | n/a | n/a | n/a | n/a | n/a | 0.05 | 1 |
| 2-Hydroxyatorvastatin   | 0.673 | 0.05 | 0.017 (40 mg) | 0.00085 | n/a | n/a | n/a | n/a | n/a | 0.05 | n/d |
| 2-Hydroxyatorvastatin lactone | 0.725 | 0.05 | 0.026 (40 mg) | 0.0013 | n/a | n/a | n/a | n/a | n/a | 0.05 | n/d |
| 4-Hydroxyatorvastatin   | 0.672 | 0.06 | n/a | n/a | n/a | n/a | n/a | n/a | n/a | 0.05 | n/d |
| 4-Hydroxyatorvastatin lactone | 0.721 | 0.05 | 0.0032 (40 mg) | 0.00016 | n/a | n/a | n/a | n/a | n/a | 0.05 | n/d |
| 3R,5S-fluvastatin       | 0.676 | 0.01 | 0.37 (40 mg) | 0.0037 | 97 | 2.9 | 0.74 | 1.9 | 0.019 | 0.05 | 1 |
| 3S,5R-fluvastatin       | 0.676 | 0.01 | 0.38 (40 mg) | 0.0038 | 97 | 2.9 | 0.74 | 1.9 | 0.019 | 0.05 | 1 |
| 2-Hydroxyatorvastatin   | 0.695 | 0.04 | 0.16 (4 mg) | 0.0064 | 19 | 0.57 | 0.53 | 0.30 | 0.012 | 0.04 | 1 |
| Pravastatin             | 0.753 | 0.05 | n/a | n/a | n/a | n/a | n/a | n/a | 0.03 | 0.01 | 1 |
| Pitavastatin lactone    | 0.662 | 0.52 | 0.079 (40 mg) | 0.041 | 380 | 11 | 11 | 3.0 | 1.6 | 0.1 | 1 |
| Rosuvastatin            | 0.674 | 0.12 | 0.036 (40 mg) | 0.0043 | 660 | 20 | 19 | 1.9 | 0.62 | 0.01 | 0.2 |
| Simvastatin (lactone)   | 0.768 | 0.06 | 0.023 (40 mg) | 0.0013 | 380 | 12 | n/a | 2.6 | 0.16 | 0.05 | 0.05 |
| Simvastatin acid        | 0.658 | 0.07 | 0.008 (40 mg lactone) | 0.00056 | n/a | n/a | n/a | n/a | n/a | 0.05 | 0.05 |

BP, blood-to-plasma concentration ratio; C₀, initial incubation concentration; C_{gut}, concentration in gut lumen; C_{max}, peak concentration in plasma; C_{max, portal}, peak concentration in portal vein; ℓ_{up}, unbound fraction in plasma; n/a, not available; n/d, not determined; u, unbound.

*Predicted value.

'C_{gut} calculated according to dose/250 ml.

'C_{max} calculated according to dose × k_{abs} × L/Q_{ent} where k_{abs} is the absorption rate constant, ℓ, is the fraction absorbed into the gut wall, and Q_{ent} is the enteroocyte blood flow (Rostami-Hodjegan and Tucker, 2004). A complete absorption scenario was assumed, using ℓ = 1, and standard values of k_{abs} = 0.03 l/min and Q_{ent} = 0.248 l/min (Obach et al., 2007; Kenny et al., 2012). Unbound concentrations were obtained by multiplying C_{ent} with unbound fraction in enterocytes (fu,ent). fu,ent values (not shown) were predicted in Simcyp Simulator v.19 (Certara UK Limited).

'C_{max, portal} calculated according to C_{max} + dose × k_{abs} × L/Q_{hep} where Q_{hep} is the hepatic blood flow (Ito et al., 1998). A complete absorption scenario was assumed, using ℓ = 1, and standard values of k_{abs} = 0.03 l/min and Q_{hep} = 20.7 ml/min/ kg (Houston and Galetin, 2008). For the calculations, the C_{max} in plasma was converted to its corresponding value in blood according to C_{max} × BP. Subsequently, the obtained C_{max, portal} in blood was converted to plasma concentration by dividing it by BP.
simvastatin, and simvastatin acid; HIMs: atorvastatin lactone and simvastatin). In
HLMs, inhibition of CYP3A4 and cytochrome P450 isoforms causing ≥50% depletions at 60 minutes in the cytochrome P450 screening were tested. The substrate
and NADPH concentrations used were identical to those used in the micro-
sonal depletions experiments above (Supplemental Table 2; Table 1). The protein concentration was 0.2 mg/ml in atorvastatin and atorvastatin lactone experiments; 0.5 mg/ml in 38,5S-fluvastatin and 35,5R-fluvastatin, pitavastatin lactone, and simvastatin acid experiments; and 0.1 mg/ml in simvastatin experiments. Ketoco-
nazole (1 μM), montelukast (5 μM), quinidine (10 μM), and sulfaphenazole (10 μM) were employed as competitive inhibitors of CYP3A4/5, CYP2C8, CYP2D6, and CYP2C9, respectively. Because of concerns regarding the selectivity
of montelukast and quinidine, the effects of the time-dependent inhibitors
gemfibrozil glucuronide (60 μM; CYP2C8) and paroxetine (15 μM; CYP2D6) were also tested. Whereas inhibitor and substrate were coincubated in direct inhibi-
tion experiments, the experiment with paroxetine included a 15-minute prein-
cubation of inhibitor and NADPH and HLMs before addition of substrate. The experiment with gemfibrozil glucuronide was initiated by preincubating inhibitor with NADPH (1 mM) in HLMs (2 mg/ml). After preincubation for 15 minutes, 10 μl of the preincubation mix was moved to another well containing 190 μl statin and NADPH (1 mM) in buffer, diluting the protein concentration 20-fold. Samples in the inhibition experiments were taken at the same time points as in the HLM depletions experiment described above or at 0.5, 2, 4, 6, 8, and 10 minutes (atorvastatin lactone). In HIM incubations, the protein concentration was 0.2 mg/ml, and only the effects of ketoconazole (1 μM) were tested. For atorva-
statin lactone, samples were taken at 0.5, 1, 2, 4, 5, 6, 8, 10, and 15 minutes and for simva-
statin at 0, 3.5, 5, 7.5, and 10 minutes. All HLM and HIM incubations were carried out once in triplicate at 37°C and 350 rpm and stopped as described above.

Determination of Unbound Fraction in Microsomes. Unbound fraction in microsomes (f_{unm}) values were measured using two-chambered rapid equilib-
rium dialysis devices (Thermo Scientific, Waltham, MA). HLMs (0.2 mg/ml; 0.1 mg/ml for simvastatin) or HIMs (0.2 mg/ml in buffer) (200 μl) containing statin were transferred to one chamber, and 400 μl buffer to the other, before incubation
at 37°C for 4 hours on a shaker (300 rpm). At the end of the incubation, 25-
μl samples from the microsomal and buffer chambers were transferred to a 96-
well plate containing 100 μl internal standard in acetone. Blank buffer or blank HLMs or HIMs (25 μl) were added to the samples from the microsomal or buffer chambers, respectively, to yield identical matrices. Samples were there-
after processed as described previously. The f_{unm} was calculated by dividing the statin concentration in buffer by that in the microsomal incubation mixture. To minimize potential metabolism by enzymes not dependent on external cofactors for their activity, old microsomes with several freeze and thaw cycles were used. In addition, they were allowed to incubate in room temperature 5 hours prior to the experiment. Denaturation was not carried out because of concerns that it might lead to conformational changes of the microsomal structure.

Inhibition of HMG-CoA. To investigate the effects of the time-dependent depletions of statins on their pharmacological target HMG-CoA reductase, we first determined the time linearity of HMG-CoA metabolism at different enzyme concentrations (0.3–1.2 µg/ml). Herein, incubations containing HMG-CoA (30 μM) and HMG-CoA reductase in phosphate buffer (0.1 M) were first preincubated for 5 minutes at 37°C before addition of NADPH (4.5 mM), which initiated the reactions. Reac-
tions were stopped by placing 50-μl samples in 150 μl ice-cold acetonitrile con-
taining mevalonolactone-d3 as the internal standard. Based on the obtained data, we determined the enzyme kinetics of HMG-CoA to its metabolite mevalonate by preincubating HMG-CoA (5–240 μM) with HMG-CoA reductase (0.9 µg/ml) in phosphate buffer for 3 minutes. NADPH was added and the reactions were allowed to continue for 3 minutes before stopping them as described above. The same preincubation and incubation times (3 + 3 minutes) were also used in the
final inhibition experiments. Herein, eight different concentrations of the statin
acids (0–300 nM) were simultaneously incubated with HMG-CoA at 20 μM and HMG-CoA reductase (0.9 µg/ml) and NADPH in buffer. All incubations were performed once in triplicate. Use of recombiant enzyme was preferred over HLMs to avoid cytochrome P450-mediated metabolism of statins.

Sample Processing and Analysis of Metabolism Samples. After sample collection, metabolism samples were kept on ice for at least 10 minutes before centrifugation at 2000g for 10 minutes. All samples were analyzed by liquid chromatography-tandem mass spectrometry (LC-MS/MS). The drug concentra-
tions in HLM and recombinant cytochrome P450 enzyme incubations were
determined using a Nexera X2 liquid chromatograph (Shimadzu, Kyoto, Japan) coupled to an APCI3000 tandem mass spectrometer (AB Sciex, Toronto, Ontario, Canada), as previously described for atorvastatin, fluvastatin, pitavastatin, pravas-
tatin, rosuvastatin, and simvastatin (Keskiotalo et al., 2009a,b,c; Ikonen et al., 2015; Lehtisalo et al., 2020). Part of the samples (HIM, HLC, rapid equilibrium
dilation, and some inhibition incubations) was analyzed using a Sciex 5500 Qtrap
LC-MS/MS system (AB Sciex) interfaced with an ESI ion source. The chro-
matographic separation was carried out on a Luna Omega polar C18 column
(100 × 2.1 mm I.D., 1.6-μm particle size; Phenomenex, Torrance, CA) using 5 mM ammonium formate (pH 3.9, adjusted with 98% formic acid) as mobile phase A and acetoneitrile as mobile phase B. The flow rate and the column tempera-
ture were maintained at 300 μl/min and 40°C. The following gradient condi-
tions were applied: 1 minute at 20% B on hold and then a linear ramp from 20% B to 40% B over 3 minutes followed by a second linear ramp to 90% B over 2
minutes and then 1 minute at 90% B before a re-equilibration step back to the
initial conditions (20% B). The characteristic multiple reaction monitoring transi-
tions for each analyte and internal standard are presented in Supplemental Table 1.

Sample Processing and Analysis of Pharmacodynamic Samples. After sample collection, samples were kept on ice, and 10 μl HCl (5 M) was added to
ensure the lactonization of mevalonic acid into mevalonolactone (Honda et al., 2007). Samples were centrifuged at 2000g for 10 minutes before LC-MS/MS
analysis. Analyses were separated on Kinetex C18, 2.6 μm, 100 × 21 mm with SecurityGuard ULTRA C18 2.1 2.6 mm (Phenomenex, Torrance, CA) by
liquid chromatography (Nexera X2, ultra high performance liquid chromatog-
raphy system; Shimadzu, Kyoto, Japan; with 0.05% formic acid as mobile phase
A and acetoneitrile as mobile phase B at flow rate 0.2 ml/min. The mobile phase
B was kept at 30% for 0–15 minutes, increased to 100% for 1.6–3.5 minutes, and
balanced at 30% before the next injection. The analytes were detected in a Shimadzu LCMS-8050 mass spectrometer (Shimadzu, Kyoto, Japan) operated in
positive electrospray ionization mode (ESI+), and the multiple reaction monitor-
ing transitions [M + H]+ > m/z 130.9-43.0 for mevalonolactone and m/z
133.9-45.95 for the internal standard.

Data Analysis and In Vitro-In Vivo Extrapolation. The obtained data of the present in vitro experiments were analyzed using GraphPad Prism software
(version 7.03; GraphPad Software, Inc., San Diego, CA). For depletation data, pseudo-first order depletions and coefficients (k_{dep}) were determined using nonlinear regression analysis (C = C_{0} × e^{−k_{dep} × t}, where C is the observed concentration, C_{0} the initial concentration, and t is the incubation time). Only data points in the log-linear portion of each depletion curve were included in the analyses. Statin depletions observed in incubations with cofactors were corrected for depletions in incubations lacking cofactors (negative controls) by k_{dep, corrected} = k_{dep, incubation} – k_{dep, negative control}. Assuming that substrate concentrations were negligible for their metabolic pathways, their intrinsic clearance in depletion experiments was expressed as CL_{int} = k_{dep, corrected}/[M], where [M] is the microsomal protein concentration or cytochrome P450 concentration in recombinant enzyme incubations (Obach, 1999). Percent inhibition of statin depletion was calculated by comparing CL_{int, valu} values of incubations containing inhibitor to those lacking inhibitor. The mean of
statin CL_{int} in the presence of each inhibitor was compared with the control using the Student’s t test in IBM SPSS Statistics (version 27.0; IBM Corp., Armonk, NY). A P value < 0.05 was considered statistically significant. Unbound intrinsic
clearance values, CL_{int, up} were calculated according to CL_{int} = CL_{int}/f_{unm}. For pharmacodynamic data, statin IC_{50} values were determined using nonlinear regres-
sion in GraphPad Prism and related to their unbound peak concentrations in plasma (C_{max, u}) and in the portal vein.

Results

Metabolism in Microsomal and Cytosolic Incubations. In HLM incubations, statin lactones were metabolized to a much higher extent than their corresponding acid forms (Figs. 1A and 2A; Supplemental Table 3). In incubations with NADPH, atorvastatin lactone and simva-
statin showed extensive metabolism (CL_{int} values of 3700 and 7400
μl/min per milligram), whereas the metabolism of the lactones of 2-hydroxyatorvastatin, 4-hydroxyatorvastatin, and pitavastatin was
slower (CL_{int} 20–840 μl/min per milligram). The statin acids had CL_{int}
values in the range of <0.1–80 μl/min per milligram. For most parent

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statins, the metabolism of both acid and lactone forms in NADPH + UDPGA incubations was of similar range or slightly lower than in NADPH incubations. No metabolism was observed in pitavastatin acid, pravastatin, and rosuvastatin incubations. In most HLM control incubations lacking cofactors (negative controls), the depletion of statins was <20% at 60 minutes (Fig. 2A). However, in control incubations containing simvastatin and 4-hydroxyatorvastatin lactone, there seemed to be some cytochrome P450- and UGT-independent depletion occurring; for simvastatin, the depletion was 29% at 60 minutes, and for 4-hydroxyatorvastatin lactone, it was 32%. Pitavastatin lactone exhibited large variation because of limited solubility, which was observed repeatedly across experiments. For most statins, the solubility seemed to be dependent on the presence of microsomal protein; the initial concentrations in buffer controls were often lower than in corresponding incubations containing microsomes, and the solubility seemed to increase with incubation time in buffer controls (data not shown). In the HLC screening, no clear statin metabolism was evident, either with or without PAPS (Fig. 2B).

In HIM incubations with NADPH and UDPGA, only atorvastatin lactone and simvastatin showed notable metabolism (Figs. 1B and 2C; Supplemental Table 4). The HIM CLₐₙₙ values of atorvastatin lactone and simvastatin corresponded to approximately 20% of those obtained in HLM incubations. After incubation for 20–25 minutes, no clear metabolism was observed in HIM incubations lacking cofactors, except for a slight 17% decrease in simvastatin acid concentration in PMSF-free HIM incubations (Fig. 2C). For atorvastatin, atorvastatin lactone, and pravastatin, the decrease was 10%–12%. For pitavastatin lactone, the concentration seemed to have increased with time. This is likely due to solubility issues, which were also observed in other incubations with pitavastatin lactone. fₐₙₙₙₙ values varied markedly between the statins (Supplemental Tables 3 and 4).

Metabolism Screening with Recombinant Cytochrome P450 Enzymes. In the recombinant cytochrome P450 screening, CYP3A4, CYP3A5, CYP2D6, CYP2C9, and CYP2C8 were the most active enzymes involved in statin metabolism (Fig. 2A). After incubation for 60 minutes, ≤50% of the initial substrate concentration remained in CYP3A4 incubations with atorvastatin lactone, 2-hydroxyatorvastatin lactone, 4-hydroxyatorvastatin lactone, pitavastatin lactone, simvastatin, and simvastatin acid. The same (<50% remaining of the initial statin concentration) was observed in CYP3A5 incubations for atorvastatin lactone, 2-hydroxyatorvastatin lactone, and simvastatin; in CYP2D6 incubations for atorvastatin lactone, 2-hydroxyatorvastatin lactone, and pitavastatin lactone; in CYP2C9 incubations for the fluvastatin enantiomers; in CYP2C8 incubations for 2-hydroxyatorvastatin lactone and simvastatin; and in CYP1A2 and CYP2A6 incubations for 2-hydroxyatorvastatin lactone. Of note, whereas the lactones of atorvastatin, 2-hydroxyatorvastatin and simvastatin, were completely metabolized by both CYP3A4 and CYP3A5, their acid forms were more prone to metabolism by CYP3A4 than by CYP3A5.

Inhibition of Metabolism in Microsomal Incubations. In HLMs, the CYP3A4 inhibitor ketoconazole completely (>95%) inhibited the metabolism (CLₐₙₙ) of atorvastatin (P < 0.001) and atorvastatin lactone (P = 0.004) and that of simvastatin (P = 0.002) and simvastatin acid (P < 0.001) (Fig. 3). It had no effect (<10% inhibition) on the depletion of 35SR-fluvastatin (P = 0.142), whereas it had a moderate effect on the CLₐₙₙ of 35SR-fluvastatin (35% inhibition, P = 0.027) and that of pitavastatin lactone (18%, P = 0.343). In turn, the CYP2C9 inhibitor sulfaphenazole inhibited the CLₐₙₙ of 3SR,5S-fluvastatin and 35SR-fluvastatin by 42% (P < 0.001) and 51% (P = 0.010). The CYP2D6 inhibitor quinidine inhibited the depletion of atorvastatin lactone by 18% (P = 0.093) and pitavastatin lactone by 18% (P = 0.241). Paroxetine inhibited the depletion of atorvastatin lactone by 16% (P = 0.106). The CYP2C8 inhibitor montelukast inhibited the depletion of simvastatin by 25% (P = 0.101), whereas gemfibrozil glucuronide had no effect on it (P = 0.840). In HIMs, ketoconazole inhibited the metabolism of atorvastatin lactone and simvastatin by >95% (P < 0.001 and P < 0.001).

Interconversion between Acid and Lactone Forms in Microsomal and Cytosolic Incubations. Some interconversion between statin acid and lactone forms could be detected for atorvastatin,

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**Fig. 1.** Intrinsic clearance values of the tested statins obtained in the depletion experiment in HLMs (A) and HIMs (B). The CLₐₙₙ values obtained in incubations fortified with either NADPH (only in HLM incubations) or NADPH + UDPGA (both HLM and HIM incubations) are shown. The protein concentration was 0.2 mg/ml in all incubations, except for simvastatin HLM incubations (0.1 mg/ml). The data represent mean and S.D. values of triplicate incubations, and they have been corrected for potential depletion in corresponding incubations lacking cofactors (negative controls). ATOR, atorvastatin; FLU, fluvastatin; lac, lactone; PITA, pitavastatin; PRA, pravastatin; ROSU, rosuvastatin; SIM, simvastatin.
pitavastatin, and simvastatin. To examine the extent of interconversion in HIMs, we quantified the concentrations of the corresponding lactone in statin acid incubations and those of the corresponding acid in lactone incubations (Supplemental Fig. 1). For other incubations, the data described reflect qualitative data.

In HIMs supplemented with NADPH + UDPGA, lactone concentrations marginally increased in pitavastatin but not in atorvastatin incubations (Supplemental Fig. 1). In HLMs with NADPH and UDPGA, the lactone concentrations of both atorvastatin and pitavastatin slightly increased, but not in other incubations (NADPH, control). For simvastatin acid, there was no increase in lactone in HLMs and HIMs. In other incubations (PMSF-free HIMs, HLC ± PAPS) with these statin acids, no increase in lactone was observed.

For statin lactones, there seemed to be a trend toward increase in evi pitavastatin and simvastatin. To examine the extent of interconversion in HIMs, we quantified the concentrations of the corresponding lactone in statin acid incubations and those of the corresponding acid in lactone incubations (Supplemental Fig. 1). For other incubations, the data described reflect qualitative data.

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Inhibition of HMG-CoA. In enzyme kinetic experiments, the kinetics of HMG-CoA was best described by a substrate inhibition model, with $K_m$, $K_i$, and $V_{max}$ values corresponding to 13 $\mu$M, 105 $\mu$M, and 1,582 nmol/min per milligram (Supplemental Fig. 2). In pharmacodynamic experiments, the majority of the statins tested inhibited the HMG-CoA-mevalonate reaction with $IC_{50}$ values in the range of 4-20 nM (Fig. 4; Table 2). The $IC_{50}$ of 3R,5S-fluvastatin was 9 nM, whereas that of its 3S,5R enantiomer approximated to 100 nM. Similarly, the $IC_{50}$ of 2-hydroxyatorvastatin acid was 12 nM, but that of 4-hydroxyatorvastatin acid corresponded to $\sim$100 nM. Although the $IC_{50}$ values of most statins were relatively close to their typical unbound peak concentrations in plasma (Fig. 4), those of 3S,5R-fluvastatin and simvastatin acid exceeded clinically relevant concentrations of 3S,5R-fluvastatin and simvastatin acid by 26- to 35-fold.

Discussion
This study aimed to comprehensively investigate and compare the metabolism and pharmacodynamics of 14 statins or statin metabolites.
the majority of the statin acids inhibited HMG-CoA reductase with IC\textsubscript{50} values from depletion studies. In agreement with previous knowledge (Fujino et al., 2004), the lactones were metabolized more extensively than their acid forms. Atorvastatin lactone and simvastatin showed extensive metabolism, with HLM CL\textsubscript{int} values of 3700 and 7400 µl/min per milligram, whereas those of the lactones of 2-hydroxyatorvastatin, 4-hydroxyatorvastatin, and pitavastatin were in the range of 20–840 µl/min per milligram. The statin acids had CL\textsubscript{int} values below 80 µl/min per milligram, with pitavastatin acid, pravastatin, and rosuvastatin showing negligible metabolism. In HIMs, only atorvastatin lactone and simvastatin exhibited notable metabolism, with CL\textsubscript{int} values corresponding to 20% of those obtained in HLMs. As compared with incubations with NADPH as the single cofactor, UDPGA addition did not result in an increased depletion. This indicates that the role of UGTs in the overall statin metabolism is relatively small.

To verify the cytochromes P450 involved in statin metabolism and to evaluate the potential role of the poorly characterized CYP2J2, we carried out an extensive screening in a panel of 11 recombinant cytochromes P450. Here, CYP3A4, CYP3A5, CYP2D6, CYP2C9, and CYP2C8 were the most active enzymes. CYP2J2 exhibited some activity toward atorvastatin lactone, 2-hydroxyatorvastatin lactone, pitavastatin lactone, simvastatin, and simvastatin acid. On the other hand, these compounds were metabolized to various degrees by almost all cytochromes P450 tested. Our inhibition data in HLMs and HIMs demonstrated a major role for CYP3A4/5 in the metabolism of atorvastatin, atorvastatin lactone, simvastatin acid, and simvastatin, in line with clinical observations (Neuvonen et al., 2006). For fluvastatin, CYP2C9 was of greater importance than CYP3A4. Furthermore, CYP2C9 was more prominent in the metabolism of 3S,5R-fluvastatin than in that of its enantiomer, as described earlier (Hirvensalo et al., 2019). Pitavastatin lactone was markedly metabolized by CYP2D6 and CYP3A4 in the screening, but its depletion was inhibited only moderately by the corresponding inhibitors in HIMs. Overall, our data demonstrating key roles for CYP3A4 and CYP2C9 in statin metabolism are line with previous literature (Fujino et al., 2004; Neuvonen et al., 2006).

The interconversion between the acid and lactone forms of statins introduces an additional level of complexity into statin pharmacokinet- ics. The process can be enzyme-mediated, spontaneous, or pH-driven and occur in both the blood stream and hepatocytes (Jemal et al., 1999; Billecke et al., 2000; Prueksaritanont et al., 2002; Hoffmann and Nowoalseki, 2008; Li et al., 2019). In line with previous studies (Prueksaritanont et al., 2002; Fujino et al., 2003; Yamada et al., 2003), we observed slight increase in lactone concentrations in some statin acid incubations with UDPGA. The underlying mechanism is likely UGT-mediated glucuronidation, followed by spontaneous lactonization (Prueksaritanont et al., 2002). For statin lactones, there was a trend toward increase of acid concentrations in negative control microsomal incubations, suggesting a role for enzymes that do not require external cofactors for their activity, such as carboxylesterases (Liederer and Borchardt, 2006). In PMSF-free HIMs, however, less than 10% of the initial lactone concentrations had been depleted at 25 minutes. For comparison, in HIMs fortified with NADPH and UDPGA, only 32% and <1% of the initial atorvastatin lactone and simvastatin (lactone) remained at 20 minutes. Thus, the role of intestinal membrane-bound esterases in lactone depletion seems to be negligible. There are contradictory findings on the role of human carboxylesterases in the conversion of simvastatin to its acid form; however, these enzymes seem to catalyze this process in blood (Vickers et al., 1990; Wang et al., 2015; Li et al., 2019). Together with SULTs, carboxylesterases are also present in smaller fractions in the cytosol (Boberg et al., 2017). In our HLC incubations (±PAPS), no statin showed a >10% reduction in their concentrations at 25 minutes. No depletion was observed for pravastatin, which has been reported to be a SULT substrate in rat liver (Kitazawa et al., 1993; Watanabe et al.,

| Compound          | IC\textsubscript{50} (µM) |
|-------------------|---------------------------|
| Atorvastatin      | 12.4 ± 4.2                |
| 2-Hydroxyatorvastatin | 12.4 ± 4.2          |
| 3R,5S-Fluvastatin | 12.4 ± 4.2                |
| 3S,5R-Fluvastatin | 12.4 ± 4.2                |
| Pitavastatin      | 12.4 ± 4.2                |
| Pravastatin       | 12.4 ± 4.2                |
| Rosuvastatin      | 12.6 ± 3.7                |
| Simvastatin       | 19.7 ± 2.0                |

*The highest concentrations tested were 100 and 500 nM.*

To test the inhibitory effects of the acid forms of the statins on their pharmacological target, eight different concentrations of the test compound were incubated with HMG-CoA (20 µM) and HMG-CoA reductase (0.9 µg/ml) for 3 minutes. The results shown describe mean values of triplicate incubations (duplicate incubations for pitavastatin). The C\textsubscript{max, u, portal} values in plasma and in the portal vein (C\textsubscript{max, u, portal}) are from Table 1. As described therein, the C\textsubscript{max, u, portal} Concentrations were calculated assuming a complete absorption scenario. ATOR, atorvastatin; FLU, fluvastatin; OH, hydroxy; PITA, pitavastatin; PRA, pravastatin; ROSU, rosuvastatin; SIM, simvastatin.

**TABLE 2**

The in vitro pharmacodynamic effects of statin acids

To test the inhibitory effects of the acid forms of the statins on their pharmacological target, eight different concentrations of the test compound were incubated with HMG-CoA (20 µM) and HMG-CoA reductase (0.9 µg/ml) for 3 minutes. The results shown describe mean values of triplicate incubations (duplicate incubations for pitavastatin). The C\textsubscript{max, u, portal} values in plasma and in the portal vein (C\textsubscript{max, u, portal}) are from Table 1. As described therein, the C\textsubscript{max, u, portal} Concentrations were calculated assuming a complete absorption scenario. ATOR, atorvastatin; FLU, fluvastatin; OH, hydroxy; PITA, pitavastatin; PRA, pravastatin; ROSU, rosuvastatin; SIM, simvastatin.
2009). As a longer incubation time might have shown some involvement of cytosolic enzymes in statin metabolism, our HLC data should be interpreted with caution.

We used substrate depletion to obtain comparable estimates of the microsomal metabolism of each statin. The depletion approach operates under the assumption that the initial substrate concentration is well below $K_{m}$ ($C_0 < K_{m}$) (Obach, 1999). In our experiments, we attempted to use equal protein concentrations (0.2 mg/mL; 0.1 mg/mL for simvastatin) and low, clinically relevant statin concentrations. For most statins, an incubation concentration of 0.05 μM was used. For pitavastatin acid and lactone and rosuvastatin the corresponding concentrations were slightly lower (0.01–0.04 μM), and for pravastatin it was higher (0.1 μM), based on initial estimations of typical unbound plasma concentrations. Regardless, in HLM incubations, the $C_0 < K_{m}$ criterion was fulfilled for all statins with $K_{m}$ values reported in the literature (Supplemental Table 2). Assuming a “worst case” scenario with complete absorption of the statin into enterocytes, our initial substrate concentrations in HMs were in general 20-fold higher than those in HLMs. For the fluvastatin enantiomers, the concentrations used did not fulfill the $C_0 < K_{m}$ criterion in HMs. However, no fluvastatin metabolism was observed in HIM incubations, and also the metabolism in HLMs was limited.

The present microsomal experiments included both buffer controls and negative controls. As most statins needed protein to dissolve in the incubations, it was not possible to distinguish between potential degradation of compound in buffer and depletion in incubations lacking cofactors. However, by correcting for depletion in negative control incubations, we were able to accurately measure the NADPH- and UDPGA-dependent microsomal metabolism. Use of microsomes instead of hepatocytes allowed us to measure metabolic $CL_{int}$ values in a system stripped from confounding factors, such as drug transporters. Interestingly, although rosuvastatin is not metabolized in microsomes, there are data showing (very slow) rosuvastatin metabolism in hepatocytes (McCormick et al., 2000). The statins are known substrates of drug transporters, in particular of OATP1B1 and breast cancer resistance protein (Pasanen et al., 2006; Keskitalo et al., 2009a; Giacomini et al., 2013). Consequently, the hepatic clearance of statins is a product of their metabolic $CL_{int}$ values in combination with their transporter $CL_{int}$ values. Together with in vitro drug transport and pharmacodynamic data in combination with clinical data, our $CL_{int}$ values can be applied in systems pharmacology modeling of statin pharmacokinetics and effects.

In the present study, we also tested the inhibitory effects of the statin acids on their pharmacological target in the liver, HMG-CoA reductase. Although previous studies have determined these effects using radiometric assays or spectrophotometry to measure NADPH consumption (Kathwala, 1991; Holdgate et al., 2003; Perchellet et al., 2009), we used LC-MS/MS to measure mevalonolactone concentrations. Overall, our findings are in good agreement with literature data (Supplemental Table 5). Our IC$_{50}$ value obtained for pravastatin (13 nM), however, is 4- to 5-fold lower than previous measurements (McTaggart et al., 2001; Perchellet et al., 2009). The low value is supported by our preliminary findings, in which the IC$_{50}$ value obtained was close to or below their typical unbound peak concentrations in plasma and in the portal vein.

Taken together, we comprehensively investigated the in vitro metabolism and pharmacodynamics of statins. Together with drug transport and clinical data, our findings are applicable for use in systems pharmacology models to prospectively predict the pharmacokinetics and pharmacological effects of statins at differentdosages.

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Authorship Contributions
Participated in research design: Fillpula, Hirvensalo, Parvianen, Niemi.
Conducted experiments: Fillpula, Hirvensalo, Parvianen, Ivaska, Lönnberg, Deng, Vinnamäki, Kurkela, Neuronen.
Performed data analysis: Fillpula, Hirvensalo, Parvianen, Ivaska.
Wrote or contributed to the writing of the manuscript: Fillpula, Hirvensalo, Parvianen, Ivaska, Lönnberg, Deng, Vinnamäki, Kurkela, Neuronen, Niemi.

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