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(54) Title: CRYSTALLINE ROSUVASTATIN INTERMEDIATE

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(54) Title: CRYSTALLINE ROSUVASTATIN INTERMEDIATE

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CRystalline ROSUVASTATIN Intermediate

RELATED APPLICATIONS

[0001] This application claims the benefit of provisional application Serial Number 60/708,920, filed August 16, 2005, and provisional application Serial Number 60/710,930, filed August 23, 2005.

FIELD OF THE INVENTION

[0002] The invention relates to a crystalline intermediate of rosuvastatin and a process for the preparation thereof.

BACKGROUND OF THE INVENTION

[0003] Rosuvastatin calcium (monocalcium bis (+) 7-[4-(4-fluorophenyl)-6-isopropyl-2-(N-methyl-N-methylsulfonilaminopyrimidin-5-yl)-(3R,5S)-dihydroxy-(E)-6-heptenoate) is an HMG-CoA reductase inhibitor, developed by Shionogi for the once daily oral treatment of hyperlipidaemia (Ann Rep, Shionogi, 1996; Direct communications, Shionogi, 8 Feb 1999 & 25 Feb 2000). Rosuvastatin calcium has the following chemical formula:

![Chemical structure of Rosuvastatin calcium]

[0004] Rosuvastatin calcium is marketed under the name CRESTOR for treatment of a mammal such as a human. According to the maker of CRESTOR, it is administered in a daily dose of from about 5mg to about 40 mg for LDL cholesterol reduction.
One of the key intermediates of the synthesis of Rosuvastatin calcium is "intermediate 21." "Intermediate 21" refers to t-butyl ester of (+)-7-(4-(4-fluorophenyl)-6-isopropyl-2-(N-methyl-N-methanesulfonylaminopyrimidin)-5-yl)-3(R)hydroxy-5-oxo-(E)-6-heptenoic acid:

![Chemical structures](image)

In USRE37,314E, the corresponding methyl ester of intermediate 21 (rather than t-butyl ester) is described as a "syrup" after column chromatography. See example 1-(4). In WO03/097614, the same intermediate having a methyl ester is described as a "thick oil." See example 2, step b. In yet another reference, WO03/087112, column chromatography is carried out to purify intermediate 21.

Generally, an oil is difficult to handle and contains impurities. Furthermore, chromatography is not preferable for use on an industrial scale.

**SUMMARY OF THE INVENTION**
[0008] One embodiment of the invention provides a crystalline rosuvastatin intermediate or an enantiomer thereof having the following structure:

![Chemical Structure](image)

wherein $R_1$ in such crystalline rosuvastatin intermediate is a carboxy protecting group.

[0009] Another embodiment of the invention provides a process for preparing the above crystalline rosuvastatin intermediate including crystallizing the intermediate from a solution having at least one organic solvent.

[0010] A further embodiment of the invention provides a process for preparing rosuvastatin, rosuvastatin lactone or a pharmaceutically acceptable salt thereof including crystallizing the rosuvastatin intermediate:

![Chemical Structure](image)

wherein $R_1$ is a carboxy protecting group, from a solution having at least one organic solvent, said organic solvent being optionally in mixture with water, and converting the crystalline intermediate to rosuvastatin, rosuvastatin lactone or a pharmaceutically acceptable salt thereof.

[0011] Another embodiment of the invention provides a pharmaceutical composition including rosuvastatin or a pharmaceutically acceptable salt thereof and at least one pharmaceutically acceptable excipient, wherein the rosuvastatin, rosuvastatin lactone or salt thereof is prepared by converting crystalline rosuvastatin intermediate having the following structure:
wherein $R_1$ is a carboxy protecting group, to rosvastatin or a pharmaceutically acceptable salt thereof.

[00012] One embodiment of the invention provides a process of preparing the above pharmaceutical composition including mixing the rosvastatin, rosvastatin lactone or a pharmaceutically acceptable salt thereof with a pharmaceutically acceptably carrier.

[00013] One embodiment of the invention provides a method of lowering LDL levels in a mammal comprising administering the pharmaceutical composition of the invention to a mammal.

**BRIEF DESCRIPTION OF THE FIGURES**

[00014] Fig 1: X-Ray Powder Diffractogram of crystalline Rosuvastatin intermediate.

[00015] Fig. 2: DSC thermogram of crystalline Rosuvastatin intermediate.

[00016] Fig. 3: FTIR spectrum of crystalline Rosuvastatin intermediate.

**DETAILED DESCRIPTION OF THE INVENTION**

[00017] One embodiment of the invention provides a crystalline intermediate ("intermediate") or an enantiomer thereof, which is used for the synthesis of rosvastatin, having the following structure:
wherein R₁ in such crystalline rosuvastatin intermediate is a carboxy protecting group.

[00018] This crystalline intermediate is suitable for use on an industrial scale, *inter alia* because crystalline forms may be easier to handle and process than oil intermediates. Crystallization also allows for purification of the intermediate.

[00019] R₁ in the crystalline rosuvastatin intermediate may be any suitable carboxy protecting group, including but not limited to phenyl. Preferably, R₁ in the crystalline rosuvastatin intermediate is a C₁ to C₄ alkyl group. In one embodiment, R₁ is a methyl group.

[00020] In a preferred embodiment, R₁ is a tert-butyl, providing “intermediate 21”:

The crystallization and isolation of intermediate 21 is illustrated in the examples.

[00021] The crystallinity of the intermediate 21 is confirmed by powder X-Ray Diffraction. Crystalline rosuvastatin intermediate 21 may be characterized by powder x-ray diffraction peaks at 10.5, 13.1, 15.4, 19.0, and 20.4 ± 0.2 degrees two theta. Crystalline rosuvastatin intermediate 21 may be further characterized by powder x-ray diffraction peaks at 11.2, 15.7, 16.6, 18.0, 18.6, 19.4, 21.8, and 23.1 ± 0.2 degrees two theta.

[00022] Crystalline rosuvastatin intermediate 21 may be characterized by an FTIR spectrum having peaks at 1543, 1380, 1153, 961, and 847 cm⁻¹. The compound
may further be characterized by an FTIR spectrum having peaks at 2980, 1606, 1508, 1440, 1340, 1223, 1100 and 1065 cm$^{-1}$.

DSC thermogram for crystalline rosuvastatin intermediate 21 shows an endothermic peak at about 100ºC, and a broad endotherm at about 220ºC.

The intermediate, including intermediate 21, may be obtained as a solid by crystallization from a solution. The solution may be that of the intermediate in one or more organic solvents, or one or more water-miscible organic solvents in a mixture with water.

Examples of suitable solvents for crystallization include C$_6$ to C$_{12}$ aromatic and C$_5$ to C$_{12}$ aliphatic hydrocarbons, C$_3$ to C$_8$ ethers, C$_3$ to C$_8$ esters, C$_3$ to C$_8$ ketones, C$_1$ to C$_5$ alcohols, C$_1$ to C$_6$ alkylamines, and C$_1$ to C$_6$ alkylethers of ethylene glycol. Specific examples of solvents include toluene, n-heptane, n-hexane, cyclohexane, cellosolve, ethyl acetate, n-butyl acetate, t-butyl acetate, methyl t-butyl ether, di-ethyl ether, tetrahydrofuran, methanol, ethanol, isopropanol, n-butanol, methyl iso-butyl ketone, diethyl carbonate, butyl lactate, acetone, acetonitrile, mixtures thereof, and mixtures of any of these water miscible organic solvents with water. An example of a water miscible solvent for use as a mixture with water is methanol.

In a typical crystallization process, the intermediate is dissolved in one of the solvents, or the mixture of the solvents as provided above. To obtain the solution, the solvent may have to be heated. Heating is preferably carried out to a temperature of about 40ºC to about 100ºC, and more preferably to a temperature of about 40ºC to about 70ºC. The solution is then preferably allowed to cool, such to a temperature of about 20ºC to about 30ºC, or room temperature. The solution may then be seeded. After seeding, the reaction mixture, which may be a slurry, may be further cooled, preferably to a temperature of about -10ºC to about 20ºC. The crystallization process may be carried out overnight, i.e., for about 8 hours.

In one embodiment, the crystallization process includes heating the solvent to a temperature of about 40ºC to about 70ºC to obtain a solution, cooling the solution to a temperature of about 20ºC to about 30ºC, seeding, cooling after seeding to a temperature of about -10ºC to about 20ºC and recovering the crystalline form.

The crystallization may result in a sticky solid, as in example 4. In such instance, such solid may be recrystallized or slurried.
Crystallization may include adding an anti-solvent to facilitate the precipitation of the intermediate. The term “anti-solvent” refers to a liquid that, when added to a solution of intermediate in a solvent, induces precipitation of intermediate. The anti-solvent may also be in a binary mixture with the solvent when the solution is prepared. Precipitation of intermediate 21 is induced by the anti-solvent when addition of the anti-solvent causes the intermediate to precipitate from the solution more rapidly or to a greater extent than the intermediate precipitates from a solution containing an equal concentration of the intermediate in the same solvent when the solution is maintained under the same conditions for the same period of time but without adding the anti-solvent. Suitable anti-solvents include water and C₅-C₁₂ cyclic or acyclic saturated hydrocarbons. Preferred anti-solvents include water, heptane, and hexane.

The resulting crystals are then recovered by conventional techniques, such as filtration. They may be washed with water or an organic solvent. The crystals are then preferably dried. The temperature may be increased or the pressure reduced to accelerate the drying process. Drying may be carried out at a temperature of about 40°C to about 100°C, under a pressure of below about 100 mmHg. Preferably, drying occurs at a temperature of about 40°C to about 60°C. Drying may also be performed under atmospheric pressure until constant weight.

The crystalline intermediate can be used to make rosuvastatin. The intermediate, which is in the form of a keto ester, is reduced to a diol ester. The reduction of the ketoester is disclosed in the art. See e.g. US2005/0159615, in regard to its processes for reduction of statins. Reagents such as RU-binap, EtB₃/NaBH₄, MeO-9-BBN/NaBH₄ and diethylmethoxyborane/NaBH₄ may be used for the reduction.

The diol ester may be further converted into a pharmaceutically acceptable salt of the statin or a lactone. For example, the diol ester obtained may be reacted with sodium or calcium hydroxide to obtain the sodium or calcium salt. It is also possible to first obtain the sodium salt by reaction with sodium hydroxide, and then convert the sodium salt to calcium salt by using a source of calcium such as calcium chloride or calcium acetate. The basic hydrolysis of the statin diol-ester may be carried out with one or more equivalents of an alkali metal or alkaline earth metal base such as NaOH or Ca(OH)₂, in organic solvents such as C₁ to C₈ ethers (tetrahydrofuran, isopropyl ether), ACN (acetonitrile), C₁ to C₅ alcohols (MeOH,
EtOH, IPA (isopropyl alcohol), propanol, butanol etc., C₃ to C₈ ketones or esters (acetone, methyl ethyl ketone, methyl isopropyl ketone, ethyl acetate). The hydrolysis may also be carried out with water, a mixture of the above solvents, or a mixture of water and the above solvents, preferably at room temperature or by heating.

[00033] The present invention comprises pharmaceutical composition comprising rosuvastatin lactone or a pharmaceutically acceptable salts, and at least one pharmaceutically acceptable excipient.

[00034] The present invention further encompasses a process for preparing a pharmaceutical formulation comprising combining rosuvastatin lactone and pharmaceutically acceptable salt with at least one pharmaceutically acceptable excipient.

[00035] The present invention further encompasses the use of rosuvastatin lactone and pharmaceutically acceptable salts for the manufacture of a pharmaceutical composition.

[00036] The compositions of rosuvastatin, preferably rosuvastatin lactone and pharmaceutically acceptable salts, more preferably rosuvastatin calcium are prepared by mixing a pharmaceutically acceptable excipient with rosuvastatin (or a pharmaceutically acceptable salt thereof), wherein said rosuvastatin is prepared from the intermediate in crystalline form.

[00037] Pharmaceutical compositions of the invention include excipients. Diluents increase the bulk of a solid pharmaceutical composition, and may make a pharmaceutical dosage form containing the composition easier for the patient and caregiver to handle. Diluents for solid compositions include, for example, microcrystalline cellulose (e.g. Avicel®), microfine cellulose, lactose, starch, pregelatinized starch, calcium carbonate, calcium sulfate, sugar, dextrates, dextrin, dextrose, dibasic calcium phosphate dihydrate, tribasic calcium phosphate, kaolin, magnesium carbonate, magnesium oxide, maltodextrin, mannitol, polymethacrylates (e.g. Eudragit®), potassium chloride, powdered cellulose, sodium chloride, sorbitol and talc.

[00038] Solid pharmaceutical compositions that are compacted into a dosage form, such as a tablet, may include excipients whose functions include helping to bind the active ingredient and other excipients together after compression. Binders for solid pharmaceutical compositions include acacia, alginic acid, carbomer (e.g.
carbopol), carboxymethylcellulose sodium, dextrin, ethyl cellulose, gelatin, guar gum, hydrogenated vegetable oil, hydroxyethyl cellulose, hydroxypropyl cellulose (e.g. Klucel®), hydroxypropyl methyl cellulose (e.g. Methocel®), liquid glucose, magnesium aluminum silicate, maltodextrin, methylcellulose, polymethacrylates, povidone (e.g. Kollidon® , Plasdone®), pregelatinized starch, sodium alginate and starch.

[00039] The dissolution rate of a compacted solid pharmaceutical composition in the patient's stomach may be increased by the addition of a disintegrant to the composition. Disintegrants include alginic acid, carboxymethylcellulose calcium, carboxymethylcellulose sodium (e.g. Ac-Di-Sol®, Primellose®), colloidal silicon dioxide, croscarmellose sodium, crospovidone (e.g. Kollidon®, Polyplasdone®), guar gum, magnesium aluminum silicate, methyl cellulose, microcrystalline cellulose, polacrilin potassium, powdered cellulose, pregelatinized starch, sodium alginate, sodium starch glycolate (e.g. Explotab®) and starch.

[00040] Glidants can be added to improve the flowability of a non-compacted solid composition and to improve the accuracy of dosing. Excipients that may function as glidants include colloidal silicon dioxide, magnesium trisilicate, powdered cellulose, starch, talc and tribasic calcium phosphate.

[00041] When a dosage form such as a tablet is made by the compaction of a powdered composition, the composition is subjected to pressure from a punch and dye. Some excipients and active ingredients have a tendency to adhere to the surfaces of the punch and dye, which can cause the product to have pitting and other surface irregularities. A lubricant can be added to the composition to reduce adhesion and ease the release of the product from the dye. Lubricants include magnesium stearate, calcium stearate, glycercyl monostearate, glycercyl palmitostearate, hydrogenated castor oil, hydrogenated vegetable oil, mineral oil, polyethylene glycol, sodium benzoate, sodium lauryl sulfate, sodium stearyl fumarate, stearic acid, talc and zinc stearate.

[00042] Flavoring agents and flavor enhancers make the dosage form more palatable to the patient. Common flavoring agents and flavor enhancers for pharmaceutical products that may be included in the composition of the present invention include maltol, vanillin, ethyl vanillin, menthol, citric acid, fumaric acid, ethyl maltol and tartaric acid.
Solid and liquid compositions may also be dyed using any pharmaceutically acceptable colorant to improve their appearance and/or facilitate patient identification of the product and unit dosage level.

In liquid pharmaceutical compositions of the invention, rosuvastatin and any other solid excipients are dissolved or suspended in a liquid carrier such as water, vegetable oil, alcohol, polyethylene glycol, propylene glycol or glycerin. Liquid pharmaceutical compositions may contain emulsifying agents to disperse uniformly throughout the composition an active ingredient or other excipient that is not soluble in the liquid carrier. Emulsifying agents that may be useful in liquid compositions of the present invention include, for example, gelatin, egg yolk, casein, cholesterol, acacia, tragacanth, chondrus, pectin, methyl cellulose, carbomer, cetostearyl alcohol and cetyl alcohol.

Liquid pharmaceutical compositions may also contain a viscosity enhancing agent to improve the mouth-feel of the product and/or coat the lining of the gastrointestinal tract. Such agents include acacia, alginic acid bentonite, carbomer, carboxymethylcellulose calcium or sodium, cetostearyl alcohol, methyl cellulose, ethylcellulose, gelatin guar gum, hydroxyethyl cellulose, hydroxypropyl cellulose, hydroxypropyl methyl cellulose, maltodextrin, polyvinyl alcohol, povidone, propylene carbonate, propylene glycol alginate, sodium alginate, sodium starch glycolate, starch tragacanth and xanthan gum.

Sweetening agents such as sorbitol, saccharin, sodium saccharin, sucrose, aspartame, fructose, mannitol and invert sugar may be added to improve the taste.

Preservatives and chelating agents such as alcohol, sodium benzoate, butylated hydroxytoluene, butylated hydroxyanisole and ethylenediamine tetraacetic acid may be added at levels safe for ingestion to improve storage stability.

According to the invention, a liquid composition may also contain a buffer such as gluconic acid, lactic acid, citric acid or acetic acid, sodium gluconate, sodium lactate, sodium citrate or sodium acetate. Selection of excipients and the amounts used may be readily determined by the formulation scientist based upon experience and consideration of standard procedures and reference works in the field.

The solid compositions of the invention include powders, granulates, aggregates and compacted compositions. The dosages include dosages suitable for oral, buccal, rectal, parenteral (including subcutaneous, intramuscular, and
intravenous), inhalant and ophthalmic administration. Although the most suitable administration in any given case will depend on the nature and severity of the condition being treated, the most preferred route of the present invention is oral. The dosages may be conveniently presented in unit dosage form and prepared by any of the methods well-known in the pharmaceutical arts.

[00050] Dosage forms include solid dosage forms like tablets, powders, capsules, suppositories, sachets, troches and losenges, as well as liquid syrups, suspensions and elixirs. The dosage form of the invention may be a capsule containing the composition, preferably a powdered or granulated solid composition of the invention, within either a hard or soft shell. The shell may be made from gelatin and optionally contain a plasticizer such as glycerin and sorbitol, and an opacifying agent or colorant.

[00051] The active ingredient and excipients may be formulated into compositions and dosage forms according to methods known in the art.

[00052] A composition for tableting or capsule filling may be prepared by wet granulation. In wet granulation, some or all of the active ingredients and excipients in powder form are blended and then further mixed in the presence of a liquid, typically water, that causes the powders to clump into granules. The granulate is screened and/or milled, dried and then screened and/or milled to the desired particle size. The granulate may then be tableted, or other excipients may be added prior to tableting, such as a glidant and/or a lubricant.

[00053] A tableting composition may be prepared conventionally by dry blending. For example, the blended composition of the actives and excipients may be compacted into a slug or a sheet and then comminuted into compacted granules. The compacted granules may subsequently be compressed into a tablet.

[00054] As an alternative to dry granulation, a blended composition may be compressed directly into a compacted dosage form using direct compression techniques. Direct compression produces a more uniform tablet without granules. Excipients that are particularly well suited for direct compression tableting include microcrystalline cellulose, spray dried lactose, dicalcium phosphate dihydrate and colloidal silica. The proper use of these and other excipients in direct compression tableting is known to those in the art with experience and skill in particular formulation challenges of direct compression tableting.
[00055] A capsule filling of the invention may comprise any of the aforementioend blends and granulates that were described with reference to tableting; however, they are not subjected to a final tableting step. The oral dosage form of the invention is preferably in the form of an oral capsule having a dosage of about 5 mg to about 40 mg, more preferably capsules of 5, 10, 20 and 40 mg.

**Solid-state Characterization**

[00056] Rosuvastatin intermediate of the invention was characterized by X-Ray powder diffraction (XRD), DSC analysis and FTIR spectroscopy.

**XRD**

[00057] XRD Diffractograms were collected on Scintag X-Ray powder diffractometer model X'TRA, Cu-tube, solid state detector. Scanning parameters: Range: 2-40 deg.20: continuous scan, Rate: 3.00 deg./min.

**Thermal analysis**

[00058] Differential Scanning Calorimetry was performed on DSC821e, Mettler Toledo.
The crucible was crimped and punched prior to analysis. Experimental Conditions: Sample weight: 3-5mg. Heating rate: 10°C/min.

**FTIR spectroscopy**

[00059] FTIR spectrum was recorded on Perkin-Elmer spectrum One Spectrometer, Diffuse Reflectance Technique.

[00060] The Sample was finely ground with Potassium bromide, and the spectrum was recorded using Potassium Bromide background in a diffused reflectance accessory.

**Examples**

"TB21" refers to the t-butyl ester of intermediate 21

**Example 1: Crystallization of TB21 in toluene**

[00061] TB21 (1.3 g, 56% assay, oil) was dissolved in toluene (1.5 ml) by heating to 60°C until homogenization. The solution was then allowed to cool to room temperature, and seeding was performed. The mixture was stirred at this temperature
overnight, not causing any precipitation. The solution was then cooled to 0°C, causing precipitation. The solid was then filtered under reduced pressure, washed with some drops of toluene and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.20 g).

**Example 2: Crystallization of TB21 in EtOAc**

[00062] TB21 (1.76 g, 56% assay, oil) was dissolved in EtOAc (1.5 ml) by heating to 60°C until homogenization. The solution was then allowed to cool to room temperature, and seeding was performed. The mixture was stirred at this temperature overnight. No precipitation was observed. The solution was then cooled to 0°C, causing precipitation. The solid was then filtered under reduced pressure, washed with some drops of EtOAc and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.35 g).

**Example 3: Crystallization of TB21 in MeOH**

[00063] TB21 (1.25 g, 56% assay, oil) was dissolved in MeOH (1.5 ml) by heating to 60°C until homogenization. The solution was then allowed to cool to room temperature, and seeding was performed. The mixture was stirred at this temperature overnight. No precipitation was observed. The solution was then cooled to 0°C, causing precipitation. The solid was then filtered under reduced pressure, washed with some drops of MeOH and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.45 g).

**Example 4: Crystallization of TB21 in MeOH:H₂O**

[00064] TB21 (20 g, 56% assay, oil) was dissolved in MeOH (20 ml) and H₂O (4 ml) at 40°C. The solution was then allowed to cool to 35°C, and seeding was performed. The mixture was allowed to cool to room temperature, and after about 30 minutes starts precipitation. After being stirred at this temperature overnight, the precipitate turned into a sticky semi-solid. The mixture was then heated to 35°C and MeOH (5 ml) was added, so the sticky solid was dissolved. The slurry was then allowed to cool to room temperature, and stirred at this temperature for 2 hours. The solid was then filtered under reduced pressure, washed few drops of MeOH:H₂O (5:1) and dried at 50°C under reduced pressure until constant weight to get solid TB21 (5.86 g).
Example 5: Crystallization of TB21 in MTBE (methyl t-Butyl ether).

[00065] TB21 (2 g, 56% assay, oil) was dissolved in MTBE (2 ml) under heating to reflux. The solution was then allowed to cool to room temperature, and seeding was performed causing precipitation. The mixture was stirred at this temperature overnight, and then cooled to 0°C for about 3 hours. The solid obtained was filtered under reduced pressure, washed with some drops of MTBE and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.48 g).

Example 6: Crystallization of TB21 in IPA

[00066] TB21 (2 g, 56% assay, oil) was dissolved in IPA (2 ml) by heating to 70°C until homogenization. The solution was then allowed to cool to room temperature, and seeding was performed. Precipitation starts about 1 hour after seeding. The mixture was stirred at room temperature overnight. The slurry was then cooled to 0°C for about 30 minutes. The solid so-obtained was filtered under reduced pressure, washed with some drops of IPA and dried at 50°C under reduced pressure for 72 hrs to get solid TB21 (0.45 g).

Example 7: Crystallization of TB21 in n-BuOH

[00067] TB21 (2 g, 56% assay, oil) was dissolved in n-BuOH (2 ml) by heating to 70°C until homogenization. The solution was then allowed to cool to room temperature, and seeding was performed. No precipitation was observed. The solution was then cooled to 0°C, causing precipitation. The slurry was stirred at this temperature for about 30 minutes. The solid was then filtered under reduced pressure, washed with few drops of n-BuOH and dried at 50°C under reduced pressure for 72 hrs to get solid TB21 (0.25 g).

Example 8: Crystallization of TB21 in MIBK (methyl-isobutyl ketone)

[00068] TB21 (2 g, 56% assay, oil) was dissolved in MIBK (2 ml) by heating to 60°C until homogenization. The solution was then allowed to cool to room temperature, and seeding was performed. No precipitation was observed. The solution was then cooled to 0°C and seeded. No precipitation was observed. The mixture was stirred at room temperature overnight, and after this time there is precipitation. The slurry was then cooled to 0°C for 2 hrs, then filtered under reduced pressure, washed
with few drops of MIBK and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.09 g).

Example 9: Crystallization of TB21 in Diethyl carbonate

[00069] TB21 (2 g, 56% assay, oil) was dissolved in DEC (2 ml) by heating to 60°C until homogenization. The solution was then allowed to cool to room temperature, and seeding was performed. No precipitation was observed. The solution was then cooled to 0°C and new seeding at this temperature induced precipitation. The slurry was stirred at room temperature overnight and then cooled to 0°C for 2 hrs. The solid so-obtained was filtered under reduced pressure, washed with few drops of DEC and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.36 g).

Example 10: Crystallization of TB21 in Butyl lactate

[00070] TB21 (2 g, 56% assay, oil) was dissolved in Butyl lactate (2 ml) at 100°C until homogenization. The solution was then allowed to cool to room temperature and seeded. No precipitation was observed. The solution was then cooled to 0°C and new seeding at this temperature did not induce precipitation. The mixture was stirred at room temperature overnight and precipitation was observed. The slurry was cooled to 0°C for 2 hrs. The solid was then filtered under reduced pressure, washed with few drops of Butyl lactate and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.20 g).

Example 11: Crystallization of TB21 in MeOH:H₂O

[00071] TB21 (2 g, 56% assay, oil) was dissolved in MeOH:H₂O (5:1, 2 ml) by heating to 55°C until homogenization. The solution was then allowed to cool to room temperature and seeded. Precipitation was observed. The mixture was stirred at room temperature overnight, and then cooled to 0°C for 2 hrs. The solid so-obtained was filtered under reduced pressure, washed with few drops of MeOH:H₂O (5:1) and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.73 g).

Example 12: Crystallization of TB21 in n-Butyl acetate

[00072] TB21 (2 g, 56% assay, oil) was dissolved in n-BuOAc (2 ml) under heating. The solution was then allowed to cool to room temperature and seeding was
performed causing precipitation. The mixture was then stirred at room temperature overnight, and then cooled to 0°C for 2 hrs. The solid so-obtained was filtered under reduced pressure, washed and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.25 g %).

Example 13: Crystallization of TB21 in IPA:H₂O
[00073] TB21 (2 g, 56% assay, oil) was dissolved in IPA (2.5 ml) and H₂O (1 ml) by heating to 55°C until homogenization. The solution was then allowed to cool to room temperature and seeding was performed. No precipitation was observed. The mixture was stirred at room temperature overnight and precipitation was observed. The slurry was then cooled to 0°C for 2 hrs. The solid so-obtained was filtered under reduced pressure, washed with few drops of IPA:H₂O (2.5:1) and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.67 g).

Example 14: Crystallization of TB21 in MeOH:H₂O
[00074] TB21 (10.68 g, 56% assay, oil) was dissolved in MeOH:H₂O (5:1, 5 ml) under heating, until homogenization. The solution was then allowed to cool to room temperature and seeding was performed. No precipitation was observed. The mixture was stirred at room temperature for 72 hours giving a thick slurry. The solid so-obtained was filtered under reduced pressure, washed with few drops of MeOH:H₂O (5:1) and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (6.33 g).

Example 15: Crystallization of TB21 in MTBE
[00075] TB21 (10 g, 56% assay, oil) was dissolved in MTBE (5 ml) by heating to reflux until homogenization. The solution was then allowed to cool to room temperature, and seeding was performed. No precipitation was observed. The mixture was stirred at room temperature for 72 hours giving a thick slurry. The solid was then filtered under reduced pressure, washed with some drops of MTBE and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (4.5 g).

Example 16: Crystallization of TB21 in Acetone:H₂O
[00076] TB21 (2 g, 56% assay, oil) was dissolved in acetone (1 ml) and H₂O (0.5 ml) by heating to 60°C until homogenization. The solution was then allowed to
cool to room temperature, and seeding was performed. No precipitation was observed. The mixture was stirred at room temperature for 18 hours. After this time, precipitation was observed. The slurry was then cooled to -10°C for 2 hours. The solid was then filtered under reduced pressure, washed with some drops of Acetone:H₂O (2:1) and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.63 g)

Example 17: Crystallization of TB21 in ACN:H₂O
[00077] TB21 (2 g, 56% assay, oil) was dissolved in ACN (1 ml) and H₂O (0.5 ml) by heating to 70°C until homogenization. The solution was then allowed to cool to room temperature, and seeding was performed. No precipitation was observed. The mixture was stirred at room temperature for 18 hours. After this time, precipitation was observed. The slurry was then cooled to -10°C for 2 hours. The solid was then filtered under reduced pressure, washed with some drops of ACN:H₂O (2:1) and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.31g)

Example 18: Crystallization of TB21 in MeOH:H₂O
[00078] TB21 (2 g, 56% assay, oil) was dissolved in MeOH:H₂O (5:1, 1 ml) by heating to 70°C until homogenization. The solution was then allowed to cool to room temperature, and seeding was performed, causing precipitation. The mixture was stirred at room temperature for 18 hours, giving a slurry. The slurry was then cooled to -10°C for 2 hours. The solid was then filtered under reduced pressure, washed with some drops of MeOH:H₂O (5:1) and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.56g)

Example 19: Crystallization of TB21 in Et₂O:MeOH
[00079] TB21 (2 g, 56% assay, oil) was suspended in Et₂O (5 ml) at 35°C. MeOH (0.5 ml) was added, causing dissolution. The solution was then allowed to cool to room temperature, and seeding was performed, not causing precipitation immediately. The solution was stirred at room temperature for 18 hours. After this time precipitation was observed. The slurry was then cooled to -10°C for 5 hours. The solid was then filtered under reduced pressure, washed with some drops of Et₂O and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.5g)
Example 20: Crystallization of TB21 in Cellosolve

[00080] TB21 (2 g, 56% assay, oil) was dissolved in Cellosolve (2 ml) by heating to 90°C until homogenization. The solution was then allowed to cool to room temperature, and seeding was performed, not causing precipitation immediately. The solution was stirred at room temperature for 18 hours. After this time precipitation was observed. The slurry was then cooled to -10°C for 5 hours. The solid was then filtered under reduced pressure, washed with some drops of Cellosolve and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.21 g)

Example 21: Crystallization of TB21 in MeOH:H₂O

[00081] TB21 (10 g, 56% assay, oil) was dissolved in a mixture MeOH:H₂O (5:1, 5 ml) by heating to 60°C until homogenization. The solution was then allowed to cool to room temperature, and seeding was performed. The mixture was stirred at room temperature for 18 hours. The solid was then filtered under reduced pressure, washed with some drops of a mixture MeOH:H₂O (5:1) and dried at 50°C under reduced pressure for 18 hrs to get solid TB21 (0.56 g)

[00082] Having thus described the invention with reference to particular preferred embodiments and illustrated it with Examples, those in the art can appreciate modifications to the invention as described and illustrated that do not depart from the spirit and scope of the invention as disclosed in the specification. The Examples are set forth to aid in understanding the invention but are not intended to, and should not be construed to, limit its scope in any way. The examples do not include detailed descriptions of conventional methods.
CLAIMS

What is claimed is:

1. A crystalline rosuvastatin intermediate or an enantiomer thereof having the following structure:

   ![Chemical Structure](image)

   wherein R₁ in such crystalline rosuvastatin intermediate is a t-butyl group.

2. The crystalline rosuvastatin intermediate of claim 1, wherein the crystalline rosuvastatin intermediate has an X-Ray Diffraction pattern with peaks at 10.5, 13.1, 15.4, 19.0, and 20.4 ±0.2 degrees two theta.

3. The crystalline rosuvastatin intermediate of claim 2, further characterized by an X-Ray Diffraction pattern with peaks at 11.2, 15.7, 16.6, 18.0, 18.6, 19.4, 21.8, and 23.1 ±0.2 degrees two theta.

4. The crystalline rosuvastatin intermediate of any one of claims 2 and 3, wherein the crystalline rosuvastatin intermediate has an FTIR spectrum with peaks at 1543, 1380, 1153, 961, and 847 cm⁻¹.

5. The crystalline rosuvastatin intermediate of claim 4, further characterized by an FTIR spectrum with peaks at: 2980, 1606, 1508, 1440, 1340, 1223, 1100, and 1065 cm⁻¹.

6. The crystalline rosuvastatin intermediate of any one of claims 2-5, wherein the crystalline rosuvastatin intermediate has a DSC thermogram with an endothermic peak at about 100°C, and a broad endotherm at about 220°C.
7. A process for preparing the crystalline rosuvastatin intermediate of any one of claims 1-6 comprising crystallizing the intermediate from a solution having at least one organic solvent.

8. The process of claim 7, wherein the organic solvent is a water miscible solvent.

9. The process of claim 8, wherein the water miscible solvent is in mixture with water.

10. The process of any one of claims 7-9, wherein crystallizing comprises heating a reaction mixture of the intermediate in the solvent to obtain a solution, followed by cooling.

11. The process of claim 10, wherein crystallizing comprises adding an anti-solvent to the solution.

12. The process of claim 11, wherein the anti-solvent is selected from the group consisting of water, heptane, and hexane.

13. The process of any one of claims 10-12, wherein crystallizing comprises seeding the solution.

14. The process of any one of claims 10-13, wherein crystallizing comprises:
   a) heating the solvent to obtain a solution;
   b) cooling;
   c) seeding; and
   d) recovering the crystalline intermediate.

15. The process of claim 14, wherein cooling is carried out before seeding.

16. The process of claim 15, further comprising cooling the solution both before and after seeding.

17. The process of any one of claims 14-16, further comprising adding an antisolvent before the recovering step.
18. The process of any one of claims 10-17, wherein heating is carried out to a temperature of about 40°C to about 100°C.

19. The process of claim 18, wherein heating is carried out to a temperature of about 40°C to about 70°C.

20. The process of any one of claims 10-19, wherein cooling is carried out to a temperature of about -10°C to about 20°C.

21. The process of any one of claims 14-20, wherein the recovered crystalline rosuvastatin intermediate is dried at a temperature of about 40°C to about 100°C.

22. The process of any one of claims 7-21, wherein the solvent is selected from the group consisting of C₆ to C₁₂ aromatic and C₅ to C₁₂ aliphatic hydrocarbons, C₃ to C₈ ethers, C₃ to C₈ esters, C₃ to C₅ ketones, and C₁ to C₅ alcohols, and mixtures thereof.

23. The process of claim 22, wherein the solvent is selected from the group consisting of toluene, n-heptane, n-hexane, cyclohexane, cellosolve, ethyl acetate, n-butyl acetate, t-butyl acetate, methyl t-butyl ether, di-ethyl ether, tetrahydrofuran, methanol, ethanol, isopropanol, n-butanol, methyl iso-butyl ketone, diethyl carbonate, butyl lactate, acetone, acetonitrile, and mixtures thereof.

24. The process of claim 9, wherein the water miscible solvent in a mixture with water is a mixture of MeOH/H₂O.

25. The process of claim 9, wherein the water miscible solvent in a mixture with water is a mixture of IPA/H₂O.

26. The process of any one of claims 7-23, wherein the solvent is toluene.

27. The process of any one of claims 7-23, wherein the solvent is acetonitrile.

28. The process of claim 9, wherein the water miscible solvent in a mixture with water is a mixture of ethanol and water.
29. The process of claim 14, wherein crystallizing comprises:
   a) heating the solvent to a temperature of about 40°C to about 70°C to obtain a solution;
   b) cooling the solution to a temperature of about 20°C to about 30°C;
   c) seeding;
   d) cooling after seeding to a temperature of about -10°C to about 20°C; and
   e) recovering the crystalline intermediate.

30. The process of any one of claims 7-29, further comprising converting the crystalline rosvastatin intermediate into a lactone or pharmaceutically acceptable salt of rosvastatin.

31. A process for preparing rosvastatin, rosvastatin lactone or a pharmaceutically acceptable salt thereof comprising crystallizing the rosvastatin intermediate:

\[
\begin{align*}
\text{F} & \quad \text{N} \\
\text{N} & \quad \text{N} \\
\text{SO}_2\text{CH}_3 & \quad \text{CO}_2\text{R}_1
\end{align*}
\]

wherein \( \text{R}_1 \) is a \( t \)-butyl group from a solution having at least one organic solvent, said organic solvent being optionally in mixture with water, and converting the crystalline intermediate to rosvastatin, rosvastatin lactone or a pharmaceutically acceptable salt thereof.

32. A process of preparing a pharmaceutical composition comprising rosvastatin, rosvastatin lactone or a pharmaceutically acceptable salt thereof and at least one pharmaceutically acceptable excipient, comprising:
   a) converting crystalline rosvastatin intermediate having the following structure:
wherein R₁ is a t-butyl group, to rosuvastatin, rosuvastatin lactone or a pharmaceutically acceptable salt thereof; and

b) mixing the rosuvastatin, rosuvastatin lactone or a pharmaceutically acceptable salt thereof with a pharmaceutically acceptably excipient.

33. Use of a process according to any one of claims 7-32, in the manufacture of rosuvastatin or a pharmaceutically acceptable salt thereof.