INDICATIONS AND USAGE

Skeletal muscle effects (e.g., myopathy and rhabdomyolysis): Risks

- Patients should be advised of the increased risk of myopathy, including rhabdomyolysis, with the 10/80-mg dose. (5.1)
- Concomitant administration of strong CYP3A4 inhibitors. (4, 5.1, 7.2)

Use with fenofibrate, gallbladder studies are indicated and alternative lipid-lowering treatments should be considered. (7.8)

- Active liver disease or unexplained persistent elevations of hepatic transaminase levels (4, 5.2)
- Women who are pregnant or may become pregnant (4, 8.1)

Concomitant administration of strong CYP3A4 inhibitors. (4, 5.1)

- Uncontrolled hypothyroidism, and renal impairment. Rare cases of rhabdomyolysis with acute renal failure secondary to myoglobinuria have been reported. (4, 5.1, 8.5, 8.6)
- Liver enzyme abnormalities: Persistent elevations in hepatic transaminases can occur. Check liver enzyme tests before initiating therapy and as clinically indicated thereafter. (5.2)

To report SUSPECTED ADVERSE REACTIONS, contact Merck Sharp & Dohme Corp., a subsidiary of Merck & Co., Inc., at 1-877-888-4231 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

DRUG INTERACTIONS

Drug Interactions Associated with Increased Risk of Myopathy/Rhabdomyolysis (2.3, 2.4, 4, 5.1, 7.1, 7.2, 7.3, 7.8, 12.3)

<table>
<thead>
<tr>
<th>Interacting Agents</th>
<th>Prescribing Recommendations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Strong CYP3A4 Inhibitors, (e.g., itraconazole, ketoconazole, posaconazole, voriconazole, erythromycin, clarithromycin, telithromycin, HIV protease inhibitors, bocepruvir, telaprevir, nefazodone, cobicistat-containing products), gemfibrozil, cyclosporine, danazol</td>
<td>Contraindicated with VYTORIN</td>
</tr>
<tr>
<td>Niacin (≥1 g/day)</td>
<td>For Chinese patients, not recommended with VYTORIN</td>
</tr>
<tr>
<td>Verapamil, diltiazem, dronedarone</td>
<td>Do not exceed 10/10 mg VYTORIN daily</td>
</tr>
<tr>
<td>Amiodarone, amlodipine, ranolazine</td>
<td>Do not exceed 10/20 mg VYTORIN daily</td>
</tr>
<tr>
<td>Lomitapide</td>
<td>For patients with HoFH, do not exceed 10/20 mg VYTORIN daily*</td>
</tr>
<tr>
<td>Daptomycin</td>
<td>Temporarily suspend VYTORIN</td>
</tr>
<tr>
<td>Grapefruit juice</td>
<td>Avoid grapefruit juice</td>
</tr>
</tbody>
</table>

* For patients with HoFH who have been taking 80 mg simvastatin chronically (e.g., for 12 months or more) without evidence of muscle toxicity, do not exceed 10/40 mg VYTORIN when taking lomitapide.

- Coumarin anticoagulants: simvastatin prolongs INR. Achieve stable INR prior to starting VYTORIN. Monitor INR frequently until stable upon initiation or alteration of VYTORIN therapy. (7.8)
- Cholestyramine: Combination decreases exposure of ezetimibe. (2.3, 7.5)
- Other Lipid-lowering Medications: Use with fenofibrate increases the risk of adverse skeletal muscle effects. Caution should be used when prescribing with VYTORIN. (5.1, 7.2)
- Fenofibrate: Combination increases exposure of ezetimibe. If cholelithiasis is suspected in a patient receiving ezetimibe and a fenofibrate, gallbladder studies are indicated and alternative lipid-lowering therapy should be considered. (7.2, 7.7, 12.3)

USE IN SPECIFIC POPULATIONS

- Moderate to severe renal impairment: Doses exceeding 10/20 mg/day should be used with caution and close monitoring (2.5, 8.6).
- Chinese patients: May be at higher risk of myopathy; monitor appropriately (5.1, 8.8).

See 17 for PATIENT COUNSELING INFORMATION and FDA-approved patient labeling.

Revised: 05/2020
1.2 Homozygous Familial Hypercholesterolemia (HoFH)
1.3 Limitations of Use

2 DOSAGE AND ADMINISTRATION
2.1 Recommended Dosing
2.2 Restricted Dosing for 10/80 mg
2.3 Co-administration with Other Drugs
2.4 Patients with Homozygous Familial Hypercholesterolemia
2.5 Patients with Renal Impairment/Chronic Kidney Disease
2.6 Geriatric Patients

3 DOSAGE FORMS AND STRENGTHS

4 CONTRAINDICATIONS

5 WARNINGS AND PRECAUTIONS
5.1 Myopathy/Rhabdomyolysis
5.2 Liver Enzymes
5.3 Endocrine Function

6 ADVERSE REACTIONS
6.1 Clinical Trials Experience
6.2 Postmarketing Experience

7 DRUG INTERACTIONS
7.1 Strong CYP3A4 Inhibitors, Cyclosporine, or Danazol
7.2 Lipid-Lowering Drugs That Can Cause Myopathy When Given Alone
7.3 Amiodarone, Dronedarone, Ranolazine, or Calcium Channel Blockers
7.4 Niacin
7.5 Cholestryramine
7.6 Digoxin
7.7 Fenofibrates (e.g., fenofibrate and fenofibric acid)
7.8 Coumarin Anticoagulants
7.9 Colchicine
7.10 Daptomycin
*Sections or subsections omitted from the full prescribing information are not listed.

8 USE IN SPECIFIC POPULATIONS

FULL PRESCRIBING INFORMATION

1 INDICATIONS AND USAGE
Therapy with lipid-altering agents should be only one component of multiple risk factor intervention in individuals at significantly increased risk for atherosclerotic vascular disease due to hypercholesterolemia. Drug therapy is indicated as an adjunct to diet when the response to a diet restricted in saturated fat and cholesterol and other nonpharmacologic measures alone has been inadequate.

1.1 Primary Hyperlipidemia
VYTORIN® is indicated for the reduction of elevated total cholesterol (total-C), low-density lipoprotein cholesterol (LDL-C), apolipoprotein B (Apo B), triglycerides (TG), and non-high-density lipoprotein cholesterol (non-HDL-C), and to increase high-density lipoprotein cholesterol (HDL-C) in patients with primary (heterozygous familial and non-familial) hyperlipidemia or mixed hyperlipidemia.

1.2 Homozygous Familial Hypercholesterolemia (HoFH)
VYTORIN is indicated for the reduction of elevated total-C and LDL-C in patients with homozygous familial hypercholesterolemia, as an adjunct to other lipid-lowering treatments (e.g., LDL apheresis) or if such treatments are unavailable.

1.3 Limitations of Use
No incremental benefit of VYTORIN on cardiovascular morbidity and mortality over and above that demonstrated for simvastatin has been established.

VYTORIN has not been studied in Fredrickson type I, III, IV, and V dyslipidemias.

2 DOSAGE AND ADMINISTRATION
2.1 Recommended Dosing
The usual dosage range is 10/10 mg/day to 10/40 mg/day. The recommended usual starting dose is 10/10 mg/day or 10/20 mg/day. VYTORIN should be taken as a single daily dose in the evening, with or without food. Patients who require a larger reduction in LDL-C (greater than 55%) may be started at 10/40 mg/day in the absence of moderate to severe renal impairment (estimated glomerular filtration rate less than 60 mL/min/1.73 m²). After initiation or titration of VYTORIN, lipid levels may be analyzed after 2 or more weeks and dosage adjusted, if needed.
2.2 Restricted Dosing for 10/80 mg

Due to the increased risk of myopathy, including rhabdomyolysis, particularly during the first year of treatment, use of the 10/80-mg dose of VYTORIN should be restricted to patients who have been taking VYTORIN 10/80 mg chronically (e.g., for 12 months or more) without evidence of muscle toxicity [see Warnings and Precautions (5.1)].

Patients who are currently tolerating the 10/80-mg dose of VYTORIN who need to be initiated on an interacting drug that is contraindicated or is associated with a dose cap for simvastatin should be switched to an alternative statin or statin-based regimen with less potential for the drug-drug interaction.

Due to the increased risk of myopathy, including rhabdomyolysis, associated with the 10/80-mg dose of VYTORIN, patients unable to achieve their LDL-C goal utilizing the 10/40-mg dose of VYTORIN should not be titrated to the 10/80-mg dose, but should be placed on alternative LDL-C-lowering treatment(s) that provides greater LDL-C lowering.

2.3 Coadministration with Other Drugs

Patients taking Verapamil, Diltiazem, or Dronedarone
- The dose of VYTORIN should not exceed 10/10 mg/day [see Warnings and Precautions (5.1), Drug Interactions (7.3), and Clinical Pharmacology (12.3)].

Patients taking Amiodarone, Amlodipine or Ranolazine
- The dose of VYTORIN should not exceed 10/20 mg/day [see Warnings and Precautions (5.1), Drug Interactions (7.3), and Clinical Pharmacology (12.3)].

Patients taking Bile Acid Sequestrants
- Dosing of VYTORIN should occur either greater than or equal to 2 hours before or greater than or equal to 4 hours after administration of a bile acid sequestrant [see Drug Interactions (7.5)].

2.4 Patients with Homozygous Familial Hypercholesterolemia

The recommended dosage for patients with homozygous familial hypercholesterolemia is VYTORIN 10/40 mg/day in the evening [see Dosage and Administration, Restricted Dosing for 10/80 mg (2.2)]. VYTORIN should be used as an adjunct to other lipid-lowering treatments (e.g., LDL apheresis) in these patients or if such treatments are unavailable.

Simvastatin exposure is approximately doubled with concomitant use of lomitapide; therefore, the dose of VYTORIN should be reduced by 50% if initiating lomitapide. VYTORIN dosage should not exceed 10/20 mg/day (or 10/40 mg/day for patients who have previously taken simvastatin 80 mg/day chronically, e.g., for 12 months or more, without evidence of muscle toxicity) while taking lomitapide.

2.5 Patients with Renal Impairment/Chronic Kidney Disease

In patients with mild renal impairment (estimated GFR greater than or equal to 60 mL/min/1.73 m²), no dosage adjustment is necessary. In patients with chronic kidney disease and estimated glomerular filtration rate less than 60 mL/min/1.73 m², the dose of VYTORIN is 10/20 mg/day in the evening. In such patients, higher doses should be used with caution and close monitoring [see Warnings and Precautions (5.1); Clinical Pharmacology (12.3)].

2.6 Geriatric Patients

No dosage adjustment is necessary in geriatric patients [see Clinical Pharmacology (12.3)].

3 DOSAGE FORMS AND STRENGTHS

- VYTORIN® 10/10, (ezetimibe 10 mg and simvastatin 10 mg tablets) are white to off-white capsule-shaped tablets with code “311” on one side.
- VYTORIN® 10/20, (ezetimibe 10 mg and simvastatin 20 mg tablets) are white to off-white capsule-shaped tablets with code “312” on one side.
- VYTORIN® 10/40, (ezetimibe 10 mg and simvastatin 40 mg tablets) are white to off-white capsule-shaped tablets with code “313” on one side.
- VYTORIN® 10/80, (ezetimibe 10 mg and simvastatin 80 mg tablets) are white to off-white capsule-shaped tablets with code “315” on one side.

4 CONTRAINDICATIONS

VYTORIN is contraindicated in the following conditions:
- Concomitant administration of strong CYP3A4 inhibitors (e.g., itraconazole, ketoconazole, posaconazole, voriconazole, HIV protease inhibitors, boceprevir, telaprevir, erythromycin,
clarithromycin, telithromycin, nefazodone, and cobicistat-containing products) [see Warnings and Precautions (5.1)].

- Concomitant administration of gemfibrozil, cyclosporine, or danazol [see Warnings and Precautions (5.1)].
- Hypersensitivity to any component of this medication [see Adverse Reactions (6.2)].
- Active liver disease or unexplained persistent elevations in hepatic transaminase levels [see Warnings and Precautions (5.2)].
- Women who are pregnant or may become pregnant. Serum cholesterol and triglycerides increase during normal pregnancy, and cholesterol or cholesterol derivatives are essential for fetal development. Because HMG-CoA reductase inhibitors (statins), such as simvastatin, decrease cholesterol synthesis and possibly the synthesis of other biologically active substances derived from cholesterol, VYTORIN may cause fetal harm when administered to a pregnant woman. Atherosclerosis is a chronic process and the discontinuation of lipid-lowering drugs during pregnancy should have little impact on the outcome of long-term therapy of primary hypercholesterolemia. There are no adequate and well-controlled studies of VYTORIN use during pregnancy; however, in rare reports congenital anomalies were observed following intrauterine exposure to statins. In rat and rabbit animal reproduction studies, simvastatin revealed no evidence of teratogenicity. VYTORIN should be administered to women of childbearing age only when such patients are highly unlikely to conceive. If the patient becomes pregnant while taking this drug, VYTORIN should be discontinued immediately and the patient should be apprised of the potential hazard to the fetus [see Use in Specific Populations (8.1)].
- Nursing mothers. It is not known whether simvastatin is excreted into human milk; however, a small amount of another drug in this class does pass into breast milk. Because statins have the potential for serious adverse reactions in nursing infants, women who require VYTORIN treatment should not breastfeed their infants [see Use in Specific Populations (8.3)].

5 WARNINGS AND PRECAUTIONS

5.1 Myopathy/Rhabdomyolysis

Simvastatin occasionally causes myopathy manifested as muscle pain, tenderness or weakness with creatine kinase above ten times the upper limit of normal (ULN). Myopathy sometimes takes the form of rhabdomyolysis with or without acute renal failure secondary to myoglobinuria, and rare fatalities have occurred. The risk of myopathy is increased by elevated plasma levels of simvastatin and simvastatin acid. Predisposing factors for myopathy include advanced age (≥65 years), female gender, uncontrolled hypothyroidism, and renal impairment. Chinese patients may be at increased risk for myopathy [see Use in Specific Populations (8.8)].

The risk of myopathy, including rhabdomyolysis, is dose related. In a clinical trial database in which 41,413 patients were treated with simvastatin, 24,747 (approximately 60%) of whom were enrolled in studies with a median follow-up of at least 4 years, the incidence of myopathy was approximately 0.03% and 0.08% at 20 and 40 mg/day, respectively. The incidence of myopathy with 80 mg (0.61%) was disproportionately higher than that observed at the lower doses. In these trials, patients were carefully monitored and some interacting medicinal products were excluded.

In a clinical trial in which 12,064 patients with a history of myocardial infarction were treated with simvastatin (mean follow-up 6.7 years), the incidence of myopathy (defined as unexplained muscle weakness or pain with a serum creatine kinase [CK] >10 times upper limit of normal [ULN]) in patients on 80 mg/day was approximately 0.9% compared with 0.02% for patients on 20 mg/day. The incidence of rhabdomyolysis (defined as myopathy with a CK >40 times ULN) in patients on 80 mg/day was approximately 0.4% compared with 0% for patients on 20 mg/day. The incidence of myopathy, including rhabdomyolysis, was highest during the first year and then notably decreased during the subsequent years of treatment. In this trial, patients were carefully monitored and some interacting medicinal products were excluded.

The risk of myopathy, including rhabdomyolysis, is greater in patients on simvastatin 80 mg compared with other statin therapies with similar or greater LDL-C-lowering efficacy and compared with lower doses of simvastatin. Therefore, the 10/80-mg dose of VYTORIN should be used only in patients who have been taking VYTORIN 10/80 mg chronically (e.g., for 12 months or more) without evidence of muscle toxicity [see Dosage and Administration, Restricted Dosing for
10/80 mg (2.2)]. If, however, a patient who is currently tolerating the 10/80-mg dose of VYTORIN needs to be initiated on an interacting drug that is contraindicated or is associated with a dose cap for simvastatin, that patient should be switched to an alternative statin or statin-based regimen with less potential for the drug-drug interaction. Patients should be advised of the increased risk of myopathy, including rhabdomyolysis, and to report promptly any unexplained muscle pain, tenderness or weakness. If symptoms occur, treatment should be discontinued immediately [see Warnings and Precautions (5.2)].

In the Study of Heart and Renal Protection (SHARP), 9270 patients with chronic kidney disease were allocated to receive VYTORIN 10/20 mg daily (n=4650) or placebo (n=4620). During a median follow-up period of 4.9 years, the incidence of myopathy (defined as unexplained muscle weakness or pain with a serum creatine kinase [CK] >10 times upper limit of normal [ULN]) was 0.2% for VYTORIN and 0.1% for placebo: the incidence of rhabdomyolysis (defined as myopathy with a CK > 40 times ULN) was 0.09% for VYTORIN and 0.02% for placebo.

In postmarketing experience with ezetimibe, cases of myopathy and rhabdomyolysis have been reported. Most patients who developed rhabdomyolysis were taking a statin prior to initiating ezetimibe. However, rhabdomyolysis has been reported with ezetimibe monotherapy and with the addition of ezetimibe to agents known to be associated with increased risk of rhabdomyolysis, such as fibric acid derivatives. VYTORIN and a fenofibrate, if taking concomitantly, should both be immediately discontinued if myopathy is diagnosed or suspected.

There have been rare reports of immune-mediated necrotizing myopathy (IMNM), an autoimmune myopathy, associated with statin use. IMNM is characterized by: proximal muscle weakness and elevated serum creatine kinase, which persist despite discontinuation of statin treatment; muscle biopsy showing necrotizing myopathy without significant inflammation; improvement with immunosuppressive agents.

All patients starting therapy with VYTORIN or whose dose of VYTORIN is being increased should be advised of the risk of myopathy, including rhabdomyolysis, and told to report promptly any unexplained muscle pain, tenderness or weakness particularly if accompanied by malaise or fever or if muscle signs and symptoms persist after discontinuing VYTORIN. VYTORIN therapy should be discontinued immediately if myopathy is diagnosed or suspected. In most cases, muscle symptoms and CK increases resolved when simvastatin treatment was promptly discontinued. Periodic CK determinations may be considered in patients starting therapy with VYTORIN or whose dose is being increased, but there is no assurance that such monitoring will prevent myopathy.

Many of the patients who have developed rhabdomyolysis on therapy with simvastatin have had complicated medical histories, including renal insufficiency usually as a consequence of long-standing diabetes mellitus. Such patients taking VYTORIN merit closer monitoring.

VYTORIN therapy should be discontinued if markedly elevated CPK levels occur or myopathy is diagnosed or suspected. VYTORIN therapy should also be temporarily withheld in any patient experiencing an acute or serious condition predisposing to the development of renal failure secondary to rhabdomyolysis, e.g., sepsis; hypotension; major surgery; trauma; severe metabolic, endocrine, or electrolyte disorders; or uncontrolled epilepsy.

**Drug Interactions**

The risk of myopathy and rhabdomyolysis is increased by elevated plasma levels of simvastatin and simvastatin acid. Simvastatin is metabolized by the cytochrome P450 isoform 3A4. Certain drugs that inhibit this metabolic pathway can raise the plasma levels of simvastatin and may increase the risk of myopathy. These include itraconazole, ketoconazole, posaconazole, and voriconazole, the macrolide antibiotics erythromycin and clarithromycin, and the ketolide antibiotic telithromycin, HIV protease inhibitors, bocaprevir, telaprevir, the antidepressant nefazodone, cobicistat-containing products, or grapefruit juice. [See Clinical Pharmacology (12.3).] Combination of these drugs with VYTORIN is contraindicated. If short-term treatment with strong CYP3A4 inhibitors is unavoidable, therapy with VYTORIN must be suspended during the course of treatment [see Contraindications (4) and Drug Interactions (7)].

The combined use of VYTORIN with gemfibrozil, cyclosporine, or danazol is contraindicated [see Contraindications (4) and Drug Interactions (7.1 and 7.2)].

Caution should be used when prescribing fenofibrates with VYTORIN, as these agents can cause myopathy when given alone and the risk is increased when they are coadministered [see Drug Interactions (7.2, 7.7)].
Cases of myopathy, including rhabdomyolysis, have been reported with simvastatin coadministered with colchicine, and caution should be exercised when prescribing VYTORIN with colchicine [see Drug Interactions (7.9)].

The benefits of the combined use of VYTORIN with the following drugs should be carefully weighed against the potential risks of combinations: other lipid-lowering drugs (fenofibrates or, for patients with HoFH, lomitapide), amiodarone, dronedarone, verapamil, diltiazem, amlodipine, or ranolazine [see Dosage and Administration (2.4), Drug Interactions (7.3)].

Cases of myopathy, including rhabdomyolysis, have been observed with simvastatin coadministered with lipid-modifying doses (≥1 g/day niacin) of niacin-containing products [see Drug Interactions (7.4)].

Cases of rhabdomyolysis have been reported with VYTORIN administered with daptomycin. Temporarily suspend VYTORIN in patients taking daptomycin [see Drug Interactions (7.10)].

Prescribing recommendations for interacting agents are summarized in Table 1 [see also Dosage and Administration (2.3, 2.4), Drug Interactions (7), and Clinical Pharmacology (12.3)].

### Table 1: Drug Interactions Associated with Increased Risk of Myopathy/Rhabdomyolysis

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<thead>
<tr>
<th>Interacting Agents</th>
<th>Prescribing Recommendations</th>
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<td>Contraindicated with VYTORIN</td>
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<td>Niacin (≥1 g/day)</td>
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<td>Verapamil, Diltiazem, Dronedarone</td>
<td>Do not exceed 10/10 mg VYTORIN daily</td>
</tr>
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<td>Amiodarone, Amlodipine, Ranolazine</td>
<td>Do not exceed 10/20 mg VYTORIN daily</td>
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<tr>
<td>Lomitapide</td>
<td>For patients with HoFH, do not exceed 10/20 mg VYTORIN daily*</td>
</tr>
<tr>
<td>Daptomycin</td>
<td>Temporarily suspend VYTORIN</td>
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<td>Grapefruit juice</td>
<td>Avoid grapefruit juice</td>
</tr>
</tbody>
</table>

*For patients with HoFH who have been taking 80 mg simvastatin chronically (e.g., for 12 months or more) without evidence of muscle toxicity, do not exceed 10/40 mg VYTORIN when taking lomitapide.

5.2 Liver Enzymes

In three placebo-controlled, 12-week trials, the incidence of consecutive elevations (≥3 X ULN) in serum transaminases was 1.7% overall for patients treated with VYTORIN and appeared to be dose-related with an incidence of 2.6% for patients treated with VYTORIN 10/80. In controlled long-term (48-week) extensions, which included both newly-treated and previously-treated patients, the incidence of consecutive elevations (≥3 X ULN) in serum transaminases was 1.8% overall and 3.6% for patients treated with VYTORIN 10/80. These elevations in transaminases were generally asymptomatic, not associated with cholestasis, and returned to baseline after discontinuation of therapy or with continued treatment.
In SHARP, 9270 patients with chronic kidney disease were allocated to receive VYTORIN 10/20 mg daily (n=4650), or placebo (n=4620). During a median follow-up period of 4.9 years, the incidence of consecutive elevations of transaminases (>3 X ULN) was 0.7% for VYTORIN and 0.6% for placebo.

It is recommended that liver function tests be performed before the initiation of treatment with VYTORIN, and thereafter when clinically indicated. There have been rare postmarketing reports of fatal and non-fatal hepatic failure in patients taking statins, including simvastatin. If serious liver injury with clinical symptoms and/or hyperbilirubinemia or jaundice occurs during treatment with VYTORIN, promptly interrupt therapy. If an alternate etiology is not found do not restart VYTORIN. Note that ALT may emanate from muscle, therefore ALT rising with CK may indicate myopathy [see Warnings and Precautions (5.1)].

VYTORIN should be used with caution in patients who consume substantial quantities of alcohol and/or have a past history of liver disease. Active liver diseases or unexplained persistent transaminase elevations are contraindications to the use of VYTORIN.

5.3 Endocrine Function

Increases in HbA1c and fasting serum glucose levels have been reported with HMG-CoA reductase inhibitors, including simvastatin.

6 ADVERSE REACTIONS

The following serious adverse reactions are discussed in greater detail in other sections of the label:

- Rhabdomyolysis and myopathy [see Warnings and Precautions (5.1)]
- Liver enzyme abnormalities [see Warnings and Precautions (5.2)]

6.1 Clinical Trials Experience

VYTORIN

Because clinical studies are conducted under widely varying conditions, adverse reaction rates observed in the clinical studies of a drug cannot be directly compared to rates in the clinical studies of another drug and may not reflect the rates observed in practice.

In the VYTORIN (ezetimibe and simvastatin) placebo-controlled clinical trials database of 1420 patients (age range 20-83 years, 52% women, 87% Caucasians, 3% Blacks, 5% Hispanics, 3% Asians) with a median treatment duration of 27 weeks, 5% of patients on VYTORIN and 2.2% of patients on placebo discontinued due to adverse reactions.

The most common adverse reactions in the group treated with VYTORIN that led to treatment discontinuation and occurred at a rate greater than placebo were:

- Increased ALT (0.9%)
- Myalgia (0.6%)
- Increased AST (0.4%)
- Back pain (0.4%)

The most commonly reported adverse reactions (incidence ≥2% and greater than placebo) in controlled clinical trials were: headache (5.8%), increased ALT (3.7%), myalgia (3.6%), upper respiratory tract infection (3.6%), and diarrhea (2.8%).

VYTORIN has been evaluated for safety in more than 10,189 patients in clinical trials.

Table 2 summarizes the frequency of clinical adverse reactions reported in ≥2% of patients treated with VYTORIN (n=1420) and at an incidence greater than placebo, regardless of causality assessment, from four placebo-controlled trials.
**Table 2**: Clinical Adverse Reactions Occurring in ≥2% of Patients Treated with VYTORIN and at an Incidence Greater than Placebo, Regardless of Causality

<table>
<thead>
<tr>
<th>Body System/Organ Class</th>
<th>Placebo (%) n=371</th>
<th>Ezetimibe 10 mg (%) n=302</th>
<th>Simvastatin† (%) n=1234</th>
<th>VYTORIN† (%) n=1420</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Body as a whole – general disorders</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Headache</td>
<td>5.4</td>
<td>6.0</td>
<td>5.9</td>
<td>5.8</td>
</tr>
<tr>
<td><strong>Gastrointestinal system disorders</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diarrhea</td>
<td>2.2</td>
<td>5.0</td>
<td>3.7</td>
<td>2.8</td>
</tr>
<tr>
<td><strong>Infections and infestations</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Influenza</td>
<td>0.8</td>
<td>1.0</td>
<td>1.9</td>
<td>2.3</td>
</tr>
<tr>
<td>Upper respiratory tract infection</td>
<td>2.7</td>
<td>5.0</td>
<td>5.0</td>
<td>3.6</td>
</tr>
<tr>
<td><strong>Musculoskeletal and connective tissue disorders</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myalgia</td>
<td>2.4</td>
<td>2.3</td>
<td>2.6</td>
<td>3.6</td>
</tr>
<tr>
<td>Pain in extremity</td>
<td>1.3</td>
<td>3.0</td>
<td>2.0</td>
<td>2.3</td>
</tr>
</tbody>
</table>

*Includes two placebo-controlled combination studies in which the active ingredients equivalent to VYTORIN were coadministered and two placebo-controlled studies in which VYTORIN was administered.
†All doses.

**Study of Heart and Renal Protection**

In SHARP, 9270 patients were allocated to VYTORIN 10/20 mg daily (n=4650) or placebo (n=4620) for a median follow-up period of 4.9 years. The proportion of patients who permanently discontinued study treatment as a result of either an adverse event or abnormal safety blood result was 10.4% vs. 9.8% among patients allocated to VYTORIN and placebo, respectively. Comparing those allocated to VYTORIN vs. placebo, the incidence of myopathy (defined as unexplained muscle weakness or pain with a serum CK >10 times ULN) was 0.2% vs. 0.1% and the incidence of rhabdomyolysis (defined as myopathy with a CK >40 times ULN) was 0.09% vs. 0.02%, respectively. Consecutive elevations of transaminases (>3 X ULN) occurred in 0.7% vs. 0.6%, respectively. Patients were asked about the occurrence of unexplained muscle pain or weakness at each study visit: 21.5% vs. 20.9% patients ever reported muscle symptoms in the VYTORIN and placebo groups, respectively. Cancer was diagnosed during the trial in 9.4% vs. 9.5% of patients assigned to VYTORIN and placebo, respectively.

**Ezetimibe**

Other adverse reactions reported with ezetimibe in placebo-controlled studies, regardless of causality assessment: **Musculoskeletal system disorders**: arthralgia; **Infections and infestations**: sinusitis; **Body as a whole – general disorders**: fatigue.

**Simvastatin**

In a clinical trial in which 12,064 patients with a history of myocardial infarction were treated with simvastatin (mean follow-up 6.7 years), the incidence of myopathy (defined as unexplained muscle weakness or pain with a serum creatine kinase [CK] >10 times upper limit of normal [ULN]) in patients on 80 mg/day was approximately 0.9% compared with 0.02% for patients on 20 mg/day. The incidence of rhabdomyolysis (defined as myopathy with a CK >40 times ULN) in patients on 80 mg/day was approximately 0.4% compared with 0% for patients on 20 mg/day. The incidence of myopathy, including rhabdomyolysis, was highest during the first year and then notably decreased during the subsequent years of treatment. In this trial, patients were carefully monitored and some interacting medicinal products were excluded.

Other adverse reactions reported with simvastatin in placebo-controlled clinical studies, regardless of causality assessment: **Cardiac disorders**: atrial fibrillation; **Ear and labyrinth disorders**: vertigo; **Gastrointestinal disorders**: abdominal pain, constipation, dyspepsia, flatulence, gastritis; **Skin and subcutaneous tissue disorders**: eczema, rash; **Endocrine disorders**: diabetes mellitus; **Infections and infestations**: bronchitis, sinusitis, urinary tract infections; **Body as a whole – general disorders**: asthenia, edema/swelling; **Psychiatric disorders**: insomnia.

**Laboratory Tests**

Marked persistent increases of hepatic serum transaminases have been noted [see Warnings and Precautions (5.2)]. Elevated alkaline phosphatase and γ-glutamyl transpeptidase have been reported. About 5% of patients taking simvastatin had elevations of CK levels of 3 or more times the normal value.
on one or more occasions. This was attributable to the noncardiac fraction of CK [see Warnings and Precautions (5.1)].

6.2 Postmarketing Experience

Because the below reactions are reported voluntarily from a population of uncertain size, it is generally not possible to reliably estimate their frequency or establish a causal relationship to drug exposure.

The following adverse reactions have been reported in postmarketing experience for VYTORIN or ezetimibe or simvastatin: pruritus; alopecia; erythema multiforme; a variety of skin changes (e.g., nodules, discoloration, dryness of skin/mucous membranes, changes to hair/nails); dizziness; muscle cramps; myalgia; arthralgia; pancreatitis; paresthesia; peripheral neuropathy; vomiting; nausea; anemia; erectile dysfunction; interstitial lung disease; myopathy/rhabdomyolysis [see Warnings and Precautions (5.1)]; hepatitis/jaundice; fatal and non-fatal hepatic failure; depression; cholelithiasis; cholecystitis; thrombocytopenia; elevations in liver transaminases; elevated creatine phosphokinase.

There have been rare reports of immune-mediated necrotizing myopathy associated with statin use [see Warnings and Precautions (5.1)].

Hypersensitivity reactions, including anaphylaxis, angioedema, rash, and urticaria have been reported. In addition, an apparent hypersensitivity syndrome has included one or more of the following features: anaphylaxis, angioedema, lupus erythematosus-like syndrome, polymyalgia rheumatica, dermatomyositis, vasculitis, purpura, thrombocytopenia, leukopenia, hemolytic anemia, positive ANA, ESR increase, eosinophilia, arthritis, arthralgia, urticaria, asthenia, photosensitivity, fever, chills, flushing, malaise, dyspnea, toxic epidermal necrolysis, erythema multiforme, including Stevens-Johnson syndrome.

There have been rare postmarketing reports of cognitive impairment (e.g., memory loss, forgetfulness, amnesia, memory impairment, confusion) associated with statin use. These cognitive issues have been reported for all statins. The reports are generally nonserious, and reversible upon statin discontinuation, with variable times to symptom onset (1 day to years) and symptom resolution (median of 3 weeks).

7 DRUG INTERACTIONS

[See Clinical Pharmacology (12.3).]

VYTORIN

7.1 Strong CYP3A4 Inhibitors, Cyclosporine, or Danazol

Strong CYP3A4 inhibitors: The risk of myopathy is increased by reducing the elimination of the simvastatin component of VYTORIN. Hence when VYTORIN is used with an inhibitor of CYP3A4 (e.g., as listed below), elevated plasma levels of HMG-CoA reductase inhibitory activity increases the risk of myopathy and rhabdomyolysis, particularly with higher doses of VYTORIN. [See Warnings and Precautions (5.1) and Clinical Pharmacology (12.3).] Concomitant use of drugs labeled as having a strong inhibitory effect on CYP3A4 is contraindicated [see Contraindications (4)]. If treatment with itraconazole, ketoconazole, posaconazole, voriconazole, erythromycin, clarithromycin or telithromycin is unavoidable, therapy with VYTORIN must be suspended during the course of treatment.

Cyclosporine or Danazol: The risk of myopathy, including rhabdomyolysis is increased by concomitant administration of cyclosporine or danazol. Therefore, concomitant use of these drugs is contraindicated [see Contraindications (4), Warnings and Precautions (5.1) and Clinical Pharmacology (12.3)].

7.2 Lipid-Lowering Drugs That Can Cause Myopathy When Given Alone

Gemfibrozil: Contraindicated with VYTORIN [see Contraindications (4) and Warnings and Precautions (5.1)].

Fenofibric acid (e.g., fenofibrate and fenofibric acid): Caution should be used when prescribing with VYTORIN [see Warnings and Precautions (5.1) and Drug Interactions (7.7)].

7.3 Amiodarone, Dronedarone, Ranolazine, or Calcium Channel Blockers

The risk of myopathy, including rhabdomyolysis, is increased by concomitant administration of amiodarone, dronedarone, ranolazine, or calcium channel blockers such as verapamil, diltiazem or amlodipine [see Dosage and Administration (2.3) and Warnings and Precautions (5.1) and Table 6 in Clinical Pharmacology (12.3)].

7.4 Niacin

Cases of myopathy/rhabdomyolysis have been observed with simvastatin coadministered with lipid-modifying doses (≥1 g/day niacin) of niacin-containing products. The risk of myopathy is greater in
Chinese patients. In a clinical trial (median follow-up 3.9 years) involving patients at high risk of cardiovascular disease and with well-controlled LDL-C levels on simvastatin 40 mg/day with or without ezetimibe 10 mg/day, there was no incremental benefit on cardiovascular outcomes with the addition of lipid-modifying doses (≥1 g/day) of niacin. Coadministration of VYTORIN with lipid-modifying doses (≥1 g/day) of niacin is not recommended in Chinese patients. It is unknown if this risk applies to other Asian patients [see Warnings and Precautions (5.1) and Use in Specific Populations (8.8)].

7.5 Cholestyramine
Concomitant cholestyramine administration decreased the mean AUC of total ezetimibe approximately 55%. The incremental LDL-C reduction due to adding VYTORIN to cholestyramine may be reduced by this interaction.

7.6 Digoxin
In one study, concomitant administration of digoxin with simvastatin resulted in a slight elevation in plasma digoxin concentrations. Patients taking digoxin should be monitored appropriately when VYTORIN is initiated.

7.7 Fenofibrates (e.g., fenofibrate and fenofibric acid)
The safety and effectiveness of VYTORIN administered with fibrates have not been established. Because it is known that the risk of myopathy during treatment with HMG-CoA reductase inhibitors is increased with concurrent administration of fenofibrates, VYTORIN should be administered with caution when used concomitantly with a fenofibrate [see Warnings and Precautions (5.1)].

Fenofibrates may increase cholesterol excretion into the bile, leading to cholelithiasis. In a preclinical study in dogs, ezetimibe increased cholesterol in the gallbladder bile [see Animal Toxicology and/or Pharmacology (13.2)]. If cholelithiasis is suspected in a patient receiving VYTORIN and a fenofibrate, gallbladder studies are indicated and alternative lipid-lowering therapy should be considered [see the product labeling for fenofibrate and fenofibric acid].

7.8 Coumarin Anticoagulants
Simvastatin 20-40 mg/day modestly potentiated the effect of coumarin anticoagulants: the prothrombin time, reported as International Normalized Ratio (INR), increased from a baseline of 1.7 to 1.8 and from 2.6 to 3.4 in a normal volunteer study and in a hypercholesterolemic patient study, respectively. With other statins, clinically evident bleeding and/or increased prothrombin time has been reported in a few patients taking coumarin anticoagulants concomitantly. In such patients, prothrombin time should be determined before starting VYTORIN and frequently enough during early therapy to ensure that no significant alteration of prothrombin time occurs. Once a stable prothrombin time has been documented, prothrombin times can be monitored at the intervals usually recommended for patients on coumarin anticoagulants. If the dose of VYTORIN is changed or discontinued, the same procedure should be repeated. Simvastatin therapy has not been associated with bleeding or with changes in prothrombin time in patients not taking anticoagulants.

Concomitant administration of ezetimibe (10 mg once daily) had no significant effect on bioavailability of warfarin and prothrombin time in a study of twelve healthy adult males. There have been postmarketing reports of increased INR in patients who had ezetimibe added to warfarin. Most of these patients were also on other medications.

The effect of VYTORIN on the prothrombin time has not been studied.

7.9 Colchicine
Cases of myopathy, including rhabdomyolysis, have been reported with simvastatin coadministered with colchicine, and caution should be exercised when prescribing VYTORIN with colchicine.

7.10 Daptomycin
Cases of rhabdomyolysis have been reported with VYTORIN administered with daptomycin. Both VYTORIN and daptomycin can cause myopathy and rhabdomyolysis when given alone and the risk of myopathy and rhabdomyolysis may be increased by coadministration. Temporarily suspend VYTORIN in patients taking daptomycin [see Warnings and Precautions (5.1)].

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy
Pregnancy Category X.
[See Contraindications (4).]
VYTORIN

VYTORIN is contraindicated in women who are or may become pregnant. Lipid-lowering drugs offer no benefit during pregnancy, because cholesterol and cholesterol derivatives are needed for normal fetal development. Atherosclerosis is a chronic process, and discontinuation of lipid-lowering drugs during pregnancy should have little impact on long-term outcomes of primary hypercholesterolemia therapy. There are no adequate and well-controlled studies of VYTORIN use during pregnancy; however, there are rare reports of congenital anomalies in infants exposed to statins in utero. Animal reproduction studies of simvastatin in rats and rabbits showed no evidence of teratogenicity. Serum cholesterol and triglycerides increase during normal pregnancy, and cholesterol or cholesterol derivatives are essential for fetal development. Because statins, such as simvastatin, decrease cholesterol synthesis and possibly the synthesis of other biologically active substances derived from cholesterol, VYTORIN may cause fetal harm when administered to a pregnant woman. If VYTORIN is used during pregnancy or if the patient becomes pregnant while taking this drug, the patient should be apprised of the potential hazard to the fetus.

Women of childbearing potential, who require VYTORIN treatment for a lipid disorder, should be advised to use effective contraception. For women trying to conceive, discontinuation of VYTORIN should be considered. If pregnancy occurs, VYTORIN should be immediately discontinued.

Ezetimibe

In oral (gavage) embryo-fetal development studies of ezetimibe conducted in rats and rabbits during organogenesis, there was no evidence of embryolethal effects at the doses tested (250, 500, 1000 mg/kg/day). In rats, increased incidences of common fetal skeletal findings (extra pair of thoracic ribs, unossified cervical vertebral centra, shortened ribs) were observed at 1000 mg/kg/day (~10 times the human exposure at 10 mg daily based on AUC0-24hr for total ezetimibe). In rabbits treated with ezetimibe, an increased incidence of extra thoracic ribs was observed at 1000 mg/kg/day (150 times the human exposure at 10 mg daily based on AUC0-24hr for total ezetimibe). Ezetimibe crossed the placenta when pregnant rats and rabbits were given multiple oral doses.

Multiple-dose studies of ezetimibe coadministered with statins in rats and rabbits during organogenesis result in higher ezetimibe and statin exposures. Reproductive findings occur at lower doses in coadministration therapy compared to monotherapy.

Simvastatin

Simvastatin was not teratogenic in rats or rabbits at doses (25, 10 mg/kg/day, respectively) that resulted in 3 times the human exposure based on mg/m² surface area. However, in studies with another structurally-related statin, skeletal malformations were observed in rats and mice.

There are rare reports of congenital anomalies following intrauterine exposure to statins. In a review¹ of approximately 100 prospectively followed pregnancies in women exposed to simvastatin or another structurally-related statin, the incidences of congenital anomalies, spontaneous abortions and fetal deaths/stillbirths did not exceed what would be expected in the general population. The number of cases is adequate only to exclude a 3- to 4-fold increase in congenital anomalies over the background incidence. In 89% of the prospectively followed pregnancies, drug treatment was initiated prior to pregnancy and was discontinued at some point in the first trimester when pregnancy was identified.

8.3 Nursing Mothers

It is not known whether simvastatin is excreted in human milk. Because a small amount of another drug in this class is excreted in human milk and because of the potential for serious adverse reactions in nursing infants, women taking simvastatin should not nurse their infants. A decision should be made whether to discontinue nursing or discontinue drug, taking into account the importance of the drug to the mother [see Contraindications (4)].

In rat studies, exposure to ezetimibe in nursing pups was up to half of that observed in maternal plasma. It is not known whether ezetimibe or simvastatin are excreted into human breast milk. Because a small amount of another drug in the same class as simvastatin is excreted in human milk and because of

the potential for serious adverse reactions in nursing infants, women who are nursing should not take VYTORIN [see Contraindications (4)].

8.4 Pediatric Use

The effects of ezetimibe coadministered with simvastatin (n=126) compared to simvastatin monotherapy (n=122) have been evaluated in adolescent boys and girls with heterozygous familial hypercholesterolemia (HeFH). In a multicenter, double-blind, controlled study followed by an open-label phase, 142 boys and 106 postmenarchal girls, 10 to 17 years of age (mean age 14.2 years, 43% females, 82% Caucasians, 4% Asian, 2% Blacks, 13% multiracial) with HeFH were randomized to receive either ezetimibe coadministered with simvastatin or simvastatin monotherapy. Inclusion in the study required 1) a baseline LDL-C level between 160 and 400 mg/dL and 2) a medical history and clinical presentation consistent with HeFH. The mean baseline LDL-C value was 225 mg/dL (range: 161-351 mg/dL) in the ezetimibe coadministered with simvastatin group compared to 219 mg/dL (range: 149-336 mg/dL) in the simvastatin monotherapy group. The patients received coadministered ezetimibe and simvastatin (10 mg, 20 mg, or 40 mg) or simvastatin monotherapy (10 mg, 20 mg, or 40 mg) for 6 weeks, coadministered ezetimibe and 40 mg simvastatin or 40 mg simvastatin monotherapy for the next 27 weeks, and open-label coadministered ezetimibe and simvastatin (10 mg, 20 mg, or 40 mg) for 20 weeks thereafter.

The results of the study at Week 6 are summarized in Table 3. Results at Week 33 were consistent with those at Week 6.

Table 3: Mean Percent Difference at Week 6 Between the Pooled Ezetimibe Coadministered with Simvastatin Group and the Pooled Simvastatin Monotherapy Group in Adolescent Patients with Heterozygous Familial Hypercholesterolemia

<table>
<thead>
<tr>
<th></th>
<th>Total-C</th>
<th>LDL-C</th>
<th>Apo B</th>
<th>Non-HDL-C</th>
<th>TG*</th>
<th>HDL-C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean percent difference</td>
<td>-12%</td>
<td>-15%</td>
<td>-12%</td>
<td>-14%</td>
<td>-2%</td>
<td>+0.1%</td>
</tr>
<tr>
<td>between treatment groups</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>95% Confidence Interval</td>
<td>(-15%, -9%)</td>
<td>(-18%, -12%)</td>
<td>(-15%, -9%)</td>
<td>(-17%, -11%)</td>
<td>(-9, +4)</td>
<td>(-3, +3)</td>
</tr>
</tbody>
</table>

* For triglycerides, median % change from baseline.

From the start of the trial to the end of Week 33, discontinuations due to an adverse reaction occurred in 7 (6%) patients in the ezetimibe coadministered with simvastatin group and in 2 (2%) patients in the simvastatin monotherapy group.

During the trial, hepatic transaminase elevations (two consecutive measurements for ALT and/or AST ≥3 X ULN) occurred in four (3%) individuals in the ezetimibe coadministered with simvastatin group and in two (2%) individuals in the simvastatin monotherapy group. Elevations of CPK (≥10 X ULN) occurred in two (2%) individuals in the ezetimibe coadministered with simvastatin group and in zero individuals in the simvastatin monotherapy group.

In this limited controlled study, there was no significant effect on growth or sexual maturation in the adolescent boys or girls, or on menstrual cycle length in girls.

Coadministration of ezetimibe with simvastatin at doses greater than 40 mg/day has not been studied in adolescents. Also, VYTORIN has not been studied in patients younger than 10 years of age or in pre-menarchal girls.

Ezetimibe

Based on total ezetimibe (ezetimibe + ezetimibe-glucuronide) there are no pharmacokinetic differences between adolescents and adults. Pharmacokinetic data in the pediatric population <10 years of age are not available.

Simvastatin

The pharmacokinetics of simvastatin has not been studied in the pediatric population.

8.5 Geriatric Use

Of the 10,189 patients who received VYTORIN in clinical studies, 3242 (32%) were 65 and older (this included 844 (8%) who were 75 and older). No overall differences in safety or effectiveness were observed between these subjects and younger subjects, and other reported clinical experience has not identified differences in responses between the elderly and younger patients but greater sensitivity of some older individuals cannot be ruled out. Since advanced age (≥65 years) is a predisposing factor for myopathy, VYTORIN should be prescribed with caution in the elderly. [See Clinical Pharmacology (12.3).]
Because advanced age (≥65 years) is a predisposing factor for myopathy, including rhabdomyolysis, VYTORIN should be prescribed with caution in the elderly. In a clinical trial of patients treated with simvastatin 80 mg/day, patients ≥65 years of age had an increased risk of myopathy, including rhabdomyolysis, compared to patients <65 years of age. [See Warnings and Precautions (5.1) and Clinical Pharmacology (12.3).]

8.6 Renal Impairment
In the SHARP trial of 9270 patients with moderate to severe renal impairment (6247 non-dialysis patients with median serum creatinine 2.5 mg/dL and median estimated glomerular filtration rate 25.6 mL/min/1.73 m\(^2\), and 3023 dialysis patients), the incidence of serious adverse events, adverse events leading to discontinuation of study treatment, or adverse events of special interest (musculoskeletal adverse events, liver enzyme abnormalities, incident cancer) was similar between patients ever assigned to VYTORIN 10/20 mg (n=4650) or placebo (n=4620) during a median follow-up of 4.9 years. However, because renal impairment is a risk factor for statin-associated myopathy, doses of VYTORIN exceeding 10/20 mg should be used with caution and close monitoring in patients with moderate to severe renal impairment. [See Dosage and Administration (2.5), Adverse Reactions (6.1), and Clinical Studies (14.3).]

8.7 Hepatic Impairment
VYTORIN is contraindicated in patients with active liver disease or unexplained persistent elevations in hepatic transaminases. [See Contraindications (4) and Warnings and Precautions (5.2).]

8.8 Chinese Patients
In a clinical trial in which patients at high risk of cardiovascular disease were treated with simvastatin 40 mg/day (median follow-up 3.9 years), the incidence of myopathy was approximately 0.05% for non-Chinese patients (n=7367) compared with 0.24% for Chinese patients (n=5468). The incidence of myopathy for Chinese patients on simvastatin 40 mg/day or ezetimibe and simvastatin 10/40 mg/day coadministered with extended-release niacin 2 g/day was 1.24%.

Chinese patients may be at higher risk for myopathy, monitor patients appropriately. Coadministration of VYTORIN with lipid-modifying doses (≥1 g/day niacin) of niacin-containing products is not recommended in Chinese patients [see Warnings and Precautions (5.1), Drug Interactions (7.4)].

10 OVERDOSAGE
VYTORIN
No specific treatment of overdosage with VYTORIN can be recommended. In the event of an overdose, symptomatic and supportive measures should be employed.

Ezetimibe
In clinical studies, administration of ezetimibe, 50 mg/day to 15 healthy subjects for up to 14 days, or 40 mg/day to 18 patients with primary hyperlipidemia for up to 56 days, was generally well tolerated.

A few cases of overdosage have been reported; most have not been associated with adverse experiences. Reported adverse experiences have not been serious.

Simvastatin
Significant lethality was observed in mice after a single oral dose of 9 g/m\(^2\). No evidence of lethality was observed in rats or dogs treated with doses of 30 and 100 g/m\(^2\), respectively. No specific diagnostic signs were observed in rodents. At these doses the only signs seen in dogs were emesis and mucoid stools.

A few cases of overdosage with simvastatin have been reported; the maximum dose taken was 3.6 g. All patients recovered without sequelae.

The dialyzability of simvastatin and its metabolites in man is not known at present.

11 DESCRIPTION
VYTORIN contains ezetimibe, a selective inhibitor of intestinal cholesterol and related phytosterol absorption, and simvastatin, an HMG-CoA reductase inhibitor.

The chemical name of ezetimibe is 1-(4-fluorophenyl)-3(R)-[3-(4-fluorophenyl)-3(S)-hydroxypropyl]-4(S)-(4-hydroxyphenyl)-2-azetidinone. The empirical formula is C\(_{24}\)H\(_{27}\)F\(_2\)NO\(_3\) and its molecular weight is 409.4.
Ezetimibe is a white, crystalline powder that is freely to very soluble in ethanol, methanol, and acetone and practically insoluble in water. Its structural formula is:

\[
\text{\begin{center}
\begin{tikzpicture}
  \node (A) at (0,0) {H};
  \node (B) at (0.5,0) {O};
  \node (C) at (1,0) {N};
  \node (D) at (1.5,0) {\text{\textbf{F}}};
  \node (E) at (2,0) {\text{\textbf{H}}};
  \node (F) at (2.5,0) {\text{\textbf{O}}};
  \node (G) at (3,0) {\text{\textbf{H}}};
  \node (H) at (3.5,0) {\text{\textbf{F}}};
  \draw (A) -- (B) -- (C) -- (D) -- (E) -- (F) -- (G) -- (H);
\end{tikzpicture}
\end{center}}
\]

Simvastatin, an inactive lactone, is hydrolyzed to the corresponding $\beta$-hydroxyacid form, which is an inhibitor of HMG-CoA reductase. Simvastatin is butanoic acid, 2,2-dimethyl-,1,2,3,7,8a-hexahydro-3,7-dimethyl-8-[2-(tetrahydro-4-hydroxy-6-oxo-2H-pyran-2-yl)-ethyl]-1-naphthalenyl ester, [1S-[1\alpha,3\alpha,7\beta,8\beta(2S^*,4S^*)]-8a\beta]]. The empirical formula of simvastatin is $\text{C}_{25}\text{H}_{38}\text{O}_{5}$ and its molecular weight is 418.57.

Simvastatin is a white to off-white, nonhygroscopic, crystalline powder that is practically insoluble in water and freely soluble in chloroform, methanol and ethanol. Its structural formula is:

\[
\text{\begin{center}
\begin{tikzpicture}
  \node (A) at (0,0) {\text{\textbf{C}}};
  \node (B) at (0.5,0) {\text{\textbf{O}}};
  \node (C) at (1,0) {\text{\textbf{O}}};
  \node (D) at (1.5,0) {\text{\textbf{H}}};
  \node (E) at (2,0) {\text{\textbf{H}}};
  \node (F) at (2.5,0) {\text{\textbf{H}}};
  \node (G) at (3,0) {\text{\textbf{H}}};
  \node (H) at (3.5,0) {\text{\textbf{H}}};
  \node (I) at (4,0) {\text{\textbf{O}}};
  \node (J) at (4.5,0) {\text{\textbf{H}}};
  \node (K) at (5,0) {\text{\textbf{H}}};
  \node (L) at (5.5,0) {\text{\textbf{H}}};
  \node (M) at (6,0) {\text{\textbf{H}}};
  \node (N) at (6.5,0) {\text{\textbf{H}}};
  \node (O) at (7,0) {\text{\textbf{H}}};
  \node (P) at (7.5,0) {\text{\textbf{H}}};
  \node (Q) at (8,0) {\text{\textbf{H}}};
  \node (R) at (8.5,0) {\text{\textbf{H}}};
  \node (S) at (9,0) {\text{\textbf{H}}};
  \node (T) at (9.5,0) {\text{\textbf{H}}};
  \node (U) at (10,0) {\text{\textbf{H}}};
  \node (V) at (10.5,0) {\text{\textbf{H}}};
  \node (W) at (11,0) {\text{\textbf{H}}};
  \node (X) at (11.5,0) {\text{\textbf{H}}};
  \node (Y) at (12,0) {\text{\textbf{H}}};
  \node (Z) at (12.5,0) {\text{\textbf{H}}};
  \node (AA) at (13,0) {\text{\textbf{H}}};
  \node (BB) at (13.5,0) {\text{\textbf{H}}};
  \node (CC) at (14,0) {\text{\textbf{H}}};
  \node (DD) at (14.5,0) {\text{\textbf{H}}};
  \node (EE) at (15,0) {\text{\textbf{H}}};
  \node (FF) at (15.5,0) {\text{\textbf{H}}};
  \node (GG) at (16,0) {\text{\textbf{H}}};
  \node (HH) at (16.5,0) {\text{\textbf{H}}};
  \node (II) at (17,0) {\text{\textbf{H}}};
  \node (JJ) at (17.5,0) {\text{\textbf{H}}};
  \node (KK) at (18,0) {\text{\textbf{H}}};
  \node (LL) at (18.5,0) {\text{\textbf{H}}};
  \node (MM) at (19,0) {\text{\textbf{H}}};
  \node (NN) at (19.5,0) {\text{\textbf{H}}};
  \node (OO) at (20,0) {\text{\textbf{H}}};
  \node (PP) at (20.5,0) {\text{\textbf{H}}};
  \node (QQ) at (21,0) {\text{\textbf{H}}};
  \node (RR) at (21.5,0) {\text{\textbf{H}}};
  \node (SS) at (22,0) {\text{\textbf{H}}};
  \node (TT) at (22.5,0) {\text{\textbf{H}}};
  \node (UU) at (23,0) {\text{\textbf{H}}};
  \node (VV) at (23.5,0) {\text{\textbf{H}}};
  \node (WW) at (24,0) {\text{\textbf{H}}};
  \node (XX) at (24.5,0) {\text{\textbf{H}}};
  \node (YY) at (25,0) {\text{\textbf{H}}};
  \node (ZZ) at (25.5,0) {\text{\textbf{H}}};
  \draw (A) -- (B) -- (C) -- (D) -- (E) -- (F) -- (G) -- (H) -- (I) -- (J) -- (K) -- (L) -- (M) -- (N) -- (O) -- (P) -- (Q) -- (R) -- (S) -- (T) -- (U) -- (V) -- (W) -- (X) -- (Y) -- (Z) -- (AA) -- (BB) -- (CC) -- (DD) -- (EE) -- (FF) -- (GG) -- (HH) -- (II) -- (JJ) -- (KK) -- (LL) -- (MM) -- (NN) -- (OO) -- (PP) -- (QQ) -- (RR) -- (SS) -- (TT) -- (UU) -- (VV) -- (WW) -- (XX) -- (YY) -- (ZZ);
\end{tikzpicture}
\end{center}}
\]

VYTORIN is available for oral use as tablets containing 10 mg of ezetimibe, and 10 mg of simvastatin (VYTORIN 10/10), 20 mg of simvastatin (VYTORIN 10/20), 40 mg of simvastatin (VYTORIN 10/40), or 80 mg of simvastatin (VYTORIN 10/80). Each tablet contains the following inactive ingredients: butylated hydroxyanisole NF, citric acid monohydrate USP, croscarmellose sodium NF, hypromellose USP, lactose monohydrate NF, magnesium stearate NF, microcrystalline cellulose NF, and propyl gallate NF.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

VYTORIN

Plasma cholesterol is derived from intestinal absorption and endogenous synthesis. VYTORIN contains ezetimibe and simvastatin, two lipid-lowering compounds with complementary mechanisms of action. VYTORIN reduces elevated total-C, LDL-C, Apo B, TG, and non-HDL-C, and increases HDL-C through dual inhibition of cholesterol absorption and synthesis.

Ezetimibe

Ezetimibe reduces blood cholesterol by inhibiting the absorption of cholesterol by the small intestine. The molecular target of ezetimibe has been shown to be the sterol transporter, Niemann-Pick C1-Like 1 (NPC1L1), which is involved in the intestinal uptake of cholesterol and phytosterols. In a 2-week clinical study in 18 hypercholesterolemic patients, ezetimibe inhibited intestinal cholesterol absorption by 54%, compared with placebo. Ezetimibe had no clinically meaningful effect on the plasma concentrations of the fat-soluble vitamins A, D, and E and did not impair adrenocortical steroid hormone production.

Ezetimibe localizes at the brush border of the small intestine and inhibits the absorption of cholesterol, leading to a decrease in the delivery of intestinal cholesterol to the liver. This causes a reduction of hepatic cholesterol stores and an increase in clearance of cholesterol from the blood; this distinct mechanism is complementary to that of statins [see Clinical Studies (14)].
Simvastatin

Simvastatin is a prodrug and is hydrolyzed to its active β-hydroxyacid form, simvastatin acid, after administration. Simvastatin is a specific inhibitor of 3-hydroxy-3-methylglutaryl-coenzyme A (HMG-CoA) reductase, the enzyme that catalyzes the conversion of HMG-CoA to mevalonate, an early and rate limiting step in the biosynthetic pathway for cholesterol. In addition, simvastatin reduces very-low-density lipoproteins (VLDL) and TG and increases HDL-C.

12.2 Pharmacodynamics

Clinical studies have demonstrated that elevated levels of total-C, LDL-C and Apo B, the major protein constituent of LDL, promote human atherosclerosis. In addition, decreased levels of HDL-C are associated with the development of atherosclerosis. Epidemiologic studies have established that cardiovascular morbidity and mortality vary directly with the level of total-C and LDL-C and inversely with the level of HDL-C. Like LDL, cholesterol-enriched triglyceride-rich lipoproteins, including VLDL, intermediate-density lipoproteins (IDL), and remnants, can also promote atherosclerosis. The independent effect of raising HDL-C or lowering TG on the risk of coronary and cardiovascular morbidity and mortality has not been determined.

12.3 Pharmacokinetics

The results of a bioequivalence study in healthy subjects demonstrated that the VYTORIN (ezetimibe and simvastatin) 10 mg/10 mg to 10 mg/80 mg combination tablets are bioequivalent to coadministration of corresponding doses of ezetimibe (ZETIA®) and simvastatin (ZOCOR®) as individual tablets.

Absorption

Ezetimibe

After oral administration, ezetimibe is absorbed and extensively conjugated to a pharmacologically active phenolic glucuronide (ezetimibe-glucuronide).

Simvastatin

The availability of the β-hydroxyacid to the systemic circulation following an oral dose of simvastatin was found to be less than 5% of the dose, consistent with extensive hepatic first-pass extraction.

Effect of Food on Oral Absorption

Ezetimibe

Concomitant food administration (high-fat or non-fat meals) had no effect on the extent of absorption of ezetimibe when administered as 10-mg tablets. The C<sub>max</sub> value of ezetimibe was increased by 38% with consumption of high-fat meals.

Simvastatin

Relative to the fasting state, the plasma profiles of both active and total inhibitors of HMG-CoA reductase were not affected when simvastatin was administered immediately before an American Heart Association recommended low-fat meal.

Distribution

Ezetimibe

Ezetimibe and ezetimibe-glucuronide are highly bound (>90%) to human plasma proteins.

Simvastatin

Both simvastatin and its β-hydroxyacid metabolite are highly bound (approximately 95%) to human plasma proteins. When radiolabeled simvastatin was administered to rats, simvastatin-derived radioactivity crossed the blood-brain barrier.

Metabolism and Excretion

Ezetimibe

Ezetimibe is primarily metabolized in the small intestine and liver via glucuronide conjugation with subsequent biliary and renal excretion. Minimal oxidative metabolism has been observed in all species evaluated.

In humans, ezetimibe is rapidly metabolized to ezetimibe-glucuronide. Ezetimibe and ezetimibe-glucuronide are the major drug-derived compounds detected in plasma, constituting approximately 10 to 20% and 80 to 90% of the total drug in plasma, respectively. Both ezetimibe and ezetimibe-glucuronide are eliminated from plasma with a half-life of approximately 22 hours for both ezetimibe and ezetimibe-glucuronide. Plasma concentration-time profiles exhibit multiple peaks, suggesting enterohepatic recycling.
Following oral administration of $^{14}$C-ezetimibe (20 mg) to human subjects, total ezetimibe (ezetimibe + ezetimibe-glucuronide) accounted for approximately 93% of the total radioactivity in plasma. After 48 hours, there were no detectable levels of radioactivity in the plasma.

Approximately 78% and 11% of the administered radioactivity were recovered in the feces and urine, respectively, over a 10-day collection period. Ezetimibe was the major component in feces and accounted for 69% of the administered dose, while ezetimibe-glucuronide was the major component in urine and accounted for 9% of the administered dose.

**Simvastatin**

Simvastatin is a lactone that is readily hydrolyzed *in vivo* to the corresponding $\beta$-hydroxyacid, a potent inhibitor of HMG-CoA reductase. Inhibition of HMG-CoA reductase is a basis for an assay in pharmacokinetic studies of the $\beta$-hydroxyacid metabolites (active inhibitors) and, following base hydrolysis, active plus latent inhibitors (total inhibitors) in plasma following administration of simvastatin. The major active metabolites of simvastatin present in human plasma are the $\beta$-hydroxyacid of simvastatin and its 6'-hydroxy, 6'-hydroxymethyl, and 6'-exomethylene derivatives.

Following an oral dose of $^{14}$C-labeled simvastatin in man, 13% of the dose was excreted in urine and 60% in feces. Plasma concentrations of total radioactivity (simvastatin plus $^{14}$C-metabolites) peaked at 4 hours and declined rapidly to about 10% of peak by 12 hours postdose.

**Specific Populations**

**Geriatric Patients**

**Ezetimibe**

In a multiple-dose study with ezetimibe given 10 mg once daily for 10 days, plasma concentrations for total ezetimibe were about 2-fold higher in older (≥65 years) healthy subjects compared to younger subjects.

**Simvastatin**

In a study including 16 elderly patients between 70 and 78 years of age who received simvastatin 40 mg/day, the mean plasma level of HMG-CoA reductase inhibitory activity was increased approximately 45% compared with 18 patients between 18-30 years of age.

**Pediatric Patients:** [See Pediatric Use (8.4).]

**Gender**

**Ezetimibe**

In a multiple-dose study with ezetimibe given 10 mg once daily for 10 days, plasma concentrations for total ezetimibe were slightly higher (<20%) in women than in men.

**Race**

**Ezetimibe**

Based on a meta-analysis of multiple-dose pharmacokinetic studies, there were no pharmacokinetic differences between Black and Caucasian subjects. Studies in Asian subjects indicated that the pharmacokinetics of ezetimibe was similar to those seen in Caucasian subjects.

**Hepatic Impairment**

**Ezetimibe**

After a single 10-mg dose of ezetimibe, the mean exposure (based on area under the curve [AUC]) to total ezetimibe was increased approximately 1.7-fold in patients with mild hepatic impairment (Child-Pugh score 5 to 6), compared to healthy subjects. The mean AUC values for total ezetimibe and ezetimibe increased approximately 3- to 4-fold and 5- to 6-fold, respectively, in patients with moderate (Child-Pugh score 7 to 9) or severe hepatic impairment (Child-Pugh score 10 to 15). In a 14-day, multiple-dose study (10 mg daily) in patients with moderate hepatic impairment, the mean AUC for total ezetimibe and ezetimibe increased approximately 4-fold compared to healthy subjects.

**Renal Impairment**

**Ezetimibe**

After a single 10-mg dose of ezetimibe in patients with severe renal disease (n=8; mean CrCl ≤30 mL/min/1.73 m²), the mean AUC for total ezetimibe and ezetimibe increased approximately 1.5-fold, compared to healthy subjects (n=9).
Simvastatin

Pharmacokinetic studies with another statin having a similar principal route of elimination to that of simvastatin have suggested that for a given dose level higher systemic exposure may be achieved in patients with severe renal impairment (as measured by creatinine clearance).

Drug Interactions [See also Drug Interactions (7).]

No clinically significant pharmacokinetic interaction was seen when ezetimibe was coadministered with simvastatin. No specific pharmacokinetic drug interaction studies with VYTORIN have been conducted other than the following study with NIASPAN (Niacin extended-release tablets).

Niacin: The effect of VYTORIN (10/20 mg daily for 7 days) on the pharmacokinetics of NIASPAN extended-release tablets (1000 mg for 2 days and 2000 mg for 5 days following a low-fat breakfast) was studied in healthy subjects. The mean C\text{max} and AUC of niacin increased 9% and 22%, respectively. The mean C\text{max} and AUC of nicotinic acid increased 10% and 19%, respectively (N=13). In the same study, the effect of NIASPAN on the pharmacokinetics of VYTORIN was evaluated (N=15). While concomitant NIASPAN decreased the mean C\text{max} of total ezetimibe (1%), and simvastatin (2%), it increased the mean C\text{max} of simvastatin acid (18%). In addition, concomitant NIASPAN increased the mean AUC of total ezetimibe (26%), simvastatin (20%), and simvastatin acid (35%).

Cases of myopathy/rhabdomyolysis have been observed with simvastatin coadministered with lipid-modifying doses (\(\geq 1\) g/day niacin) of niacin-containing products. [See Warnings and Precautions (5.1) and Drug Interactions (7.4).]

Cytochrome P450: Ezetimibe had no significant effect on a series of probe drugs (caffeine, dextromethorphan, tolbutamide, and IV midazolam) known to be metabolized by cytochrome P450 (1A2, 2D6, 2C8/9 and 3A4) in a “cocktail” study of twelve healthy adult males. This indicates that ezetimibe is neither an inhibitor nor an inducer of these cytochrome P450 isozymes, and it is unlikely that ezetimibe will affect the metabolism of drugs that are metabolized by these enzymes.

In a study of 12 healthy volunteers, simvastatin at the 80-mg dose had no effect on the metabolism of the probe cytochrome P450 isofrom 3A4 (CYP3A4) substrates midazolam and erythromycin. This indicates that simvastatin is not an inhibitor of CYP3A4 and, therefore, is not expected to affect the plasma levels of other drugs metabolized by CYP3A4.

Simvastatin acid is a substrate of the transport protein OATP1B1. Concomitant administration of medicinal products that are inhibitors of the transport protein OATP1B1 may lead to increased plasma concentrations of simvastatin acid and an increased risk of myopathy. For example, cyclosporine has been shown to increase the AUC of statins; although the mechanism is not fully understood, the increase in AUC for simvastatin acid is presumably due, in part, to inhibition of CYP3A4 and/or OATP1B1.

Simvastatin is a substrate for CYP3A4. Inhibitors of CYP3A4 can raise the plasma levels of HMG-CoA reductase inhibitory activity and increase the risk of myopathy. [See Warnings and Precautions (5.1); Drug Interactions (7.1).]
Ezetimibe

Table 4: Effect of Coadministered Drugs on Total Ezetimibe

<table>
<thead>
<tr>
<th>Coadministered Drug and Dosing Regimen</th>
<th>Total Ezetimibe*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Change in AUC</td>
</tr>
<tr>
<td>Cyclosporine-stable dose required (75-150 mg BID)&lt;sup&gt;†&lt;/sup&gt;</td>
<td>↑240%</td>
</tr>
<tr>
<td>Fenofibrate, 200 mg QD, 14 days&lt;sup&gt;‡&lt;/sup&gt;</td>
<td>↑48%</td>
</tr>
<tr>
<td>Gemfibrozil, 600 mg BID, 7 days&lt;sup&gt;‡&lt;/sup&gt;</td>
<td>↑64%</td>
</tr>
<tr>
<td>Cholestyramine, 4 g BID, 14 days&lt;sup&gt;‡&lt;/sup&gt;</td>
<td>↓55%</td>
</tr>
<tr>
<td>Aluminum &amp; magnesium hydroxide combination antacid, single dose&lt;sup&gt;§&lt;/sup&gt;</td>
<td>↓4%</td>
</tr>
<tr>
<td>Cimetidine, 400 mg BID, 7 days</td>
<td>↑16%</td>
</tr>
<tr>
<td>Glipizide, 10 mg, single dose</td>
<td>↑14%</td>
</tr>
</tbody>
</table>

Statins

<table>
<thead>
<tr>
<th>Coadministered Drug and Dosing Regimen</th>
<th>Total Ezetimibe*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Change in AUC</td>
</tr>
<tr>
<td>Lovastatin 20 mg QD, 7 days</td>
<td>↑19%</td>
</tr>
<tr>
<td>Pravastatin 20 mg QD, 14 days</td>
<td>↑7%</td>
</tr>
<tr>
<td>Atorvastatin 10 mg QD, 14 days</td>
<td>↑2%</td>
</tr>
<tr>
<td>Rosuvastatin 10 mg QD, 14 days</td>
<td>↑13%</td>
</tr>
<tr>
<td>Fluvastatin 20 mg QD, 14 days</td>
<td>↓19%</td>
</tr>
</tbody>
</table>

* Based on 10 mg-dose of ezetimibe.
† Post-renal transplant patients with mild impaired or normal renal function. In a different study, a renal transplant patient with severe renal insufficiency (creatinine clearance of 13.2 mL/min/1.73 m<sup>2</sup>) who was receiving multiple medications, including cyclosporine, demonstrated a 12-fold greater exposure to total ezetimibe compared to healthy subjects.
‡ See 7. Drug Interactions.
§ Supralox, 20 mL.

Table 5: Effect of Ezetimibe Coadministration on Systemic Exposure to Other Drugs

<table>
<thead>
<tr>
<th>Coadministered Drug and its Dosage Regimen</th>
<th>Ezetimibe Dosage Regimen</th>
<th>Change in AUC of Coadministered Drug</th>
<th>Change in C&lt;sub&gt;max&lt;/sub&gt; of Coadministered Drug</th>
</tr>
</thead>
<tbody>
<tr>
<td>Warfarin, 25 mg single dose on Day 7</td>
<td>10 mg QD, 11 days</td>
<td>↓2% (R-warfarin)</td>
<td>↑3% (R-warfarin)</td>
</tr>
<tr>
<td>Digoxin, 0.5 mg single dose</td>
<td>10 mg QD, 8 days</td>
<td>↓2%</td>
<td>↓7%</td>
</tr>
<tr>
<td>Gemfibrozil, 600 mg BID, 7 days&lt;sup&gt;‡&lt;/sup&gt;</td>
<td>10 mg QD, 7 days</td>
<td>↓1%</td>
<td>↓17%</td>
</tr>
<tr>
<td>Ethinyl estradiol &amp; Levonorgestrel, QD, 21 days</td>
<td>10 mg QD, Days 8-14 of 21 day oral contraceptive cycle</td>
<td>Ethinyl estradiol 0% Levonorgestrel 0%</td>
<td>Ethinyl estradiol 9% Levonorgestrel 5%</td>
</tr>
<tr>
<td>Glipizide, 10 mg on Days 1 and 9</td>
<td>10 mg QD, Days 2-9</td>
<td>↓3%</td>
<td>↓5%</td>
</tr>
<tr>
<td>Fenofibrate, 200 mg QD, 14 days&lt;sup&gt;‡&lt;/sup&gt;</td>
<td>10 mg QD, 14 days</td>
<td>↑111%</td>
<td>↑17%</td>
</tr>
<tr>
<td>Cyclosporine, 100 mg single dose Day 7&lt;sup&gt;‡&lt;/sup&gt;</td>
<td>20 mg QD, 8 days</td>
<td>↑15%</td>
<td>↑10%</td>
</tr>
</tbody>
</table>

Statins

<table>
<thead>
<tr>
<th>Coadministered Drug and Dosing Regimen</th>
<th>Ezetimibe Dosage Regimen</th>
<th>Change in AUC of Coadministered Drug</th>
<th>Change in C&lt;sub&gt;max&lt;/sub&gt; of Coadministered Drug</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lovastatin 20 mg QD, 7 days</td>
<td>10 mg QD, 7 days</td>
<td>↑19%</td>
<td>↑3%</td>
</tr>
<tr>
<td>Pravastatin 20 mg QD, 14 days</td>
<td>10 mg QD, 14 days</td>
<td>↓1%</td>
<td>↓24%</td>
</tr>
<tr>
<td>Atorvastatin 10 mg QD, 14 days</td>
<td>10 mg QD, 14 days</td>
<td>↓4%</td>
<td>↑7%</td>
</tr>
<tr>
<td>Rosuvastatin 10 mg QD, 14 days</td>
<td>10 mg QD, 14 days</td>
<td>↑19%</td>
<td>↑17%</td>
</tr>
<tr>
<td>Fluvastatin 20 mg QD, 14 days</td>
<td>10 mg QD, 14 days</td>
<td>↓39%</td>
<td>↓27%</td>
</tr>
</tbody>
</table>

* See 7. Drug Interactions.
### Table 6: Effect of Coadministered Drugs or Grapefruit Juice on Simvastatin Systemic Exposure

<table>
<thead>
<tr>
<th>Coadministered Drug or Grapefruit Juice</th>
<th>Dosing of Coadministered Drug or Grapefruit Juice</th>
<th>Dosing of Simvastatin</th>
<th>Geometric Mean Ratio (Ratio with / without coadministered drug)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>AUC</td>
</tr>
<tr>
<td><strong>Contraindicated with VYTORIN [see Contraindications (4) and Warnings and Precautions (5.1)]</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Telithromycin†</td>
<td>200 mg QD for 4 days</td>
<td>80 mg simvastatin ‡</td>
<td>12</td>
</tr>
<tr>
<td>Nelfinavir†</td>
<td>1250 mg BID for 14 days</td>
<td>20 mg QD for 28 days simvastatin ‡</td>
<td>6</td>
</tr>
<tr>
<td>Itraconazole†</td>
<td>200 mg QD for 4 days</td>
<td>80 mg simvastatin ‡</td>
<td>13.1</td>
</tr>
<tr>
<td>Posaconazole</td>
<td>100 mg (oral suspension) QD for 13 days</td>
<td>40 mg simvastatin ‡</td>
<td>2.85</td>
</tr>
<tr>
<td>Gemfibrozil</td>
<td>600 mg BID for 3 days</td>
<td>40 mg simvastatin ‡</td>
<td>1.35</td>
</tr>
<tr>
<td><strong>Avoid grapefruit juice with VYTORIN [see Warnings and Precautions (5.1)]</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Grapefruit Juice§ (high dose)</td>
<td>200 mL of double-strength TID¶</td>
<td>60 mg single dose simvastatin</td>
<td>7</td>
</tr>
<tr>
<td>Grapefruit Juice§ (low dose)</td>
<td>8 oz (about 237 mL) of single-strength§ #</td>
<td>20 mg single dose simvastatin</td>
<td>1.3</td>
</tr>
<tr>
<td><strong>Avoid taking with &gt;10/10 mg VYTORIN, based on clinical and/or postmarketing simvastatin experience [see Warnings and Precautions (5.1)]</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Verapamil SR</td>
<td>240 mg QD Days 1-7 then 240 mg BID on Days 8-10</td>
<td>80 mg on Day 10 simvastatin</td>
<td>2.3</td>
</tr>
<tr>
<td>Diltiazem</td>
<td>120 mg BID for 10 days</td>
<td>80 mg on Day 10 simvastatin</td>
<td>2.69</td>
</tr>
<tr>
<td>Diltiazem</td>
<td>120 mg BID for 14 days</td>
<td>20 mg on Day 14 simvastatin</td>
<td>4.8</td>
</tr>
<tr>
<td>Dronedarone</td>
<td>400 mg BID for 14 days</td>
<td>40 mg QD for 14 days simvastatin</td>
<td>1.96</td>
</tr>
<tr>
<td><strong>Avoid taking with &gt;10/20 mg VYTORIN, based on clinical and/or postmarketing simvastatin experience [see Warnings and Precautions (5.1)]</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Amiodarone</td>
<td>400 mg QD for 3 days</td>
<td>40 mg on Day 3 simvastatin</td>
<td>1.75</td>
</tr>
<tr>
<td>Amlodipine</td>
<td>10 mg QD for 10 days</td>
<td>80 mg on Day 10 simvastatin</td>
<td>1.58</td>
</tr>
<tr>
<td>Ranolazine SR</td>
<td>1000 mg BID for 7 days</td>
<td>80 mg on Day 1 and Days 6-9 simvastatin</td>
<td>2.26</td>
</tr>
</tbody>
</table>
Avoid taking with >10/20 mg VYTORIN (or 10/40 mg for patients who have previously taken 80 mg simvastatin chronically, e.g., for 12 months or more, without evidence of muscle toxicity), based on clinical experience

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dose</th>
<th>Procedure</th>
<th>Simvastatin acid</th>
<th>Simvastatin</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>mg QD for 7 days</td>
<td>40 mg single dose</td>
<td>1.7</td>
<td>1.6</td>
</tr>
<tr>
<td></td>
<td>10 mg QD for 7 days</td>
<td>20 mg single dose</td>
<td>1.4</td>
<td>1.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1.6</td>
<td>1.7</td>
</tr>
</tbody>
</table>

No dosing adjustments required for the following:

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dose</th>
<th>Procedure</th>
<th>Simvastatin acid</th>
<th>Simvastatin</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>160 mg QD for 14 days</td>
<td>80 mg QD on Days 8-14</td>
<td>0.64</td>
<td>0.89</td>
</tr>
<tr>
<td></td>
<td>80 mg single dose</td>
<td>total inhibitor</td>
<td>0.79</td>
<td>↓ from 33.6 to 21.1 ng·eq/mL</td>
</tr>
<tr>
<td></td>
<td></td>
<td>active inhibitor</td>
<td>0.79</td>
<td>↓ from 7.0 to 4.7 ng·eq/mL</td>
</tr>
</tbody>
</table>

* Results based on a chemical assay except results with propranolol as indicated.
† Results could be representative of the following CYP3A4 inhibitors: ketoconazole, erythromycin, clarithromycin, HIV protease inhibitors, and nefazodone.
‡ Simvastatin acid refers to the β-hydroxyacid of simvastatin.
§ The effect of amounts of grapefruit juice between those used in these two studies on simvastatin pharmacokinetics has not been studied.
¶ Double-strength: one can of frozen concentrate diluted with one can of water. Grapefruit juice was administered TID for 2 days, and 200 mL together with single dose simvastatin and 30 and 90 minutes following single dose simvastatin on Day 3.
* Single-strength: one can of frozen concentrate diluted with 3 cans of water. Grapefruit juice was administered with breakfast for 3 days, and simvastatin was administered in the evening on Day 3.

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

VYTORIN

No animal carcinogenicity or fertility studies have been conducted with the combination of ezetimibe and simvastatin. The combination of ezetimibe with simvastatin did not show evidence of mutagenicity in vitro in a microbial mutagenicity (Ames) test with *Salmonella typhimurium* and *Escherichia coli* with or without metabolic activation. No evidence of clastogenicity was observed in vitro in a chromosomal aberration assay in human peripheral blood lymphocytes with ezetimibe and simvastatin with or without metabolic activation. There was no evidence of genotoxicity at doses up to 600 mg/kg with the combination of ezetimibe and simvastatin (1:1) in the in vivo mouse micronucleus test.

**Ezetimibe**

A 104-week dietary carcinogenicity study with ezetimibe was conducted in rats at doses up to 1500 mg/kg/day (males) and 500 mg/kg/day (females) (~20 times the human exposure at 10 mg daily based on AUC$_{0-24 hr}$ for total ezetimibe). A 104-week dietary carcinogenicity study with ezetimibe was also conducted in mice at doses up to 500 mg/kg/day (~150 times the human exposure at 10 mg daily based on AUC$_{0-24 hr}$ for total ezetimibe). There were no statistically significant increases in tumor incidences in drug-treated rats or mice.

No evidence of mutagenicity was observed in vitro in a microbial mutagenicity (Ames) test with *Salmonella typhimurium* and *Escherichia coli* with or without metabolic activation. No evidence of clastogenicity was observed in vitro in a chromosomal aberration assay in human peripheral blood lymphocytes with or without metabolic activation. In addition, there was no evidence of genotoxicity in the in vivo mouse micronucleus test.

In oral (gavage) fertility studies of ezetimibe conducted in rats, there was no evidence of reproductive toxicity at doses up to 1000 mg/kg/day in male or female rats (~7 times the human exposure at 10 mg daily based on AUC$_{0-24 hr}$ for total ezetimibe).

**Simvastatin**

In a 72-week carcinogenicity study, mice were administered daily doses of simvastatin of 25, 100, and 400 mg/kg body weight, which resulted in mean plasma drug levels approximately 1, 4, and 8 times higher than the mean human plasma drug level, respectively, (as total inhibitory activity based on AUC)
after an 80-mg oral dose. Liver carcinomas were significantly increased in high-dose females and mid- and high-dose males with a maximum incidence of 90% in males. The incidence of adenomas of the liver was significantly increased in mid- and high-dose females. Drug treatment also significantly increased the incidence of lung adenomas in mid- and high-dose males and females. Adenomas of the Harderian gland (a gland of the eye of rodents) were significantly higher in high-dose mice than in controls. No evidence of a tumorigenic effect was observed at 25 mg/kg/day.

In a separate 92-week carcinogenicity study in mice at doses up to 25 mg/kg/day, no evidence of a tumorigenic effect was observed (mean plasma drug levels were 1 times higher than humans given 80 mg simvastatin as measured by AUC).

In a two-year study in rats at 25 mg/kg/day, there was a statistically significant increase in the incidence of thyroid follicular adenomas in female rats exposed to approximately 11 times higher levels of simvastatin than in humans given 80 mg simvastatin (as measured by AUC).

A second two-year rat carcinogenicity study with doses of 50 and 100 mg/kg/day produced hepatocellular adenomas and carcinomas (in female rats at both doses and in males at 100 mg/kg/day). Thyroid follicular cell adenomas were increased in males and females at both doses; thyroid follicular cell carcinomas were increased in females at 100 mg/kg/day. The increased incidence of thyroid neoplasms appears to be consistent with findings from other statins. These treatment levels represented plasma drug levels (AUC) of approximately 7 and 15 times (males) and 22 and 25 times (females) the mean human plasma drug exposure after an 80-mg daily dose.

No evidence of mutagenicity was observed in a microbial mutagenicity (Ames) test with or without rat or mouse liver metabolic activation. In addition, no evidence of damage to genetic material was noted in an in vitro alkaline elution assay using rat hepatocytes, a V-79 mammalian cell forward mutation study, an in vitro chromosome aberration study in CHO cells, or an in vivo chromosomal aberration assay in mouse bone marrow.

There was decreased fertility in male rats treated with simvastatin for 34 weeks at 25 mg/kg body weight (4 times the maximum human exposure level, based on AUC, in patients receiving 80 mg/day); however, this effect was not observed during a subsequent fertility study in which simvastatin was administered at this same dose level to male rats for 11 weeks (the entire cycle of spermatogenesis including epididymal maturation). No microscopic changes were observed in the testes of rats from either study. At 180 mg/kg/day (which produces exposure levels 22 times higher than those in humans taking 80 mg/day based on surface area, mg/m²), seminiferous tubule degeneration (necrosis and loss of spermatogenic epithelium) was observed. In dogs, there was drug-related testicular atrophy, decreased spermatogenesis, spermatocytic degeneration and giant cell formation at 10 mg/kg/day (approximately 2 times the human exposure, based on AUC, at 80 mg/day). The clinical significance of these findings is unclear.

13.2 Animal Toxicology and/or Pharmacology

CNS Toxicity

Optic nerve degeneration was seen in clinically normal dogs treated with simvastatin for 14 weeks at 180 mg/kg/day, a dose that produced mean plasma drug levels about 12 times higher than the mean plasma drug level in humans taking 80 mg/day.

A chemically similar drug in this class also produced optic nerve degeneration (Wallerian degeneration of retinogeniculate fibers) in clinically normal dogs in a dose-dependent fashion starting at 60 mg/kg/day, a dose that produced mean plasma drug levels about 30 times higher than the mean plasma drug level in humans taking the highest recommended dose (as measured by total enzyme inhibitory activity). This same drug also produced vestibulocochlear Wallerian-like degeneration and retinal ganglion cell chromatolysis in dogs treated for 14 weeks at 180 mg/kg/day, a dose that resulted in a mean plasma drug level similar to that seen with the 60 mg/kg/day dose.

CNS vascular lesions, characterized by perivascular hemorrhage and edema, mononuclear cell infiltration of perivascular spaces, perivascular fibrin deposits and necrosis of small vessels, were seen in dogs treated with simvastatin at a dose of 360 mg/kg/day, a dose that produced mean plasma drug levels that were about 14 times higher than the mean plasma drug levels in humans taking 80 mg/day. Similar CNS vascular lesions have been observed with several other drugs of this class.

There were cataracts in female rats after two years of treatment with 50 and 100 mg/kg/day (22 and 25 times the human AUC at 80 mg/day, respectively) and in dogs after three months at 90 mg/kg/day (19 times) and at two years at 50 mg/kg/day (5 times).
Ezetimibe

The hypocholesterolemic effect of ezetimibe was evaluated in cholesterol-fed Rhesus monkeys, dogs, rats, and mouse models of human cholesterol metabolism. Ezetimibe was found to have an ED\textsubscript{50} value of 0.5 $\mu$g/kg/day for inhibiting the rise in plasma cholesterol levels in monkeys. The ED\textsubscript{50} values in dogs, rats, and mice were 7, 30, and 700 $\mu$g/kg/day, respectively. These results are consistent with ezetimibe being a potent cholesterol absorption inhibitor.

In a rat model, where the glucuronide metabolite of ezetimibe (ezetimibe-glucuronide) was administered intraduodenally, the metabolite was as potent as ezetimibe in inhibiting the absorption of cholesterol, suggesting that the glucuronide metabolite had activity similar to the parent drug.

In 1-month studies in dogs given ezetimibe (0.03 to 300 mg/kg/day), the concentration of cholesterol in gallbladder bile increased ~2- to 4-fold. However, a dose of 300 mg/kg/day administered to dogs for one year did not result in gallstone formation or any other adverse hepatobiliary effects. In a 14-day study in mice given ezetimibe (0.3 to 5 mg/kg/day) and fed a low-fat or cholesterol-rich diet, the concentration of cholesterol in gallbladder bile was either unaffected or reduced to normal levels, respectively.

A series of acute preclinical studies was performed to determine the selectivity of ezetimibe for inhibiting cholesterol absorption. Ezetimibe inhibited the absorption of \textsuperscript{14}C-cholesterol with no effect on the absorption of triglycerides, fatty acids, bile acids, progesterone, ethinyl estradiol, or the fat-soluble vitamins A and D.

In 4- to 12-week toxicity studies in mice, ezetimibe did not induce cytochrome P450 drug-metabolizing enzymes. In toxicity studies, a pharmacokinetic interaction of ezetimibe with statins (parents or their active hydroxy acid metabolites) was seen in rats, dogs, and rabbits.

14    CLINICAL STUDIES

14.1 Primary Hyperlipidemia

VYTORIN

VYTORIN reduces total-C, LDL-C, Apo B, TG, and non-HDL-C, and increases HDL-C in patients with hyperlipidemia. Maximal to near maximal response is generally achieved within 2 weeks and maintained during chronic therapy.

VYTORIN is effective in men and women with hyperlipidemia. Experience in non-Caucasians is limited and does not permit a precise estimate of the magnitude of the effects of VYTORIN.

Five multicenter, double-blind studies conducted with either VYTORIN or coadministered ezetimibe and simvastatin equivalent to VYTORIN in patients with primary hyperlipidemia are reported: two were comparisons with simvastatin, two were comparisons with atorvastatin, and one was a comparison with rosvastatin.

In a multicenter, double-blind, placebo-controlled, 12-week trial, 1528 hyperlipidemic patients were randomized to one of ten treatment groups: placebo, ezetimibe (10 mg), simvastatin (10 mg, 20 mg, 40 mg, or 80 mg), or VYTORIN (10/10, 10/20, 10/40, or 10/80).

When patients receiving VYTORIN were compared to those receiving all doses of simvastatin, VYTORIN significantly lowered total-C, LDL-C, Apo B, TG, and non-HDL-C. The effects of VYTORIN on HDL-C were similar to the effects seen with simvastatin. Further analysis showed VYTORIN significantly increased HDL-C compared with placebo. (See Table 7.) The lipid response to VYTORIN was similar in patients with TG levels greater than or less than 200 mg/dL.
Table 7: Response to VYTORIN in Patients with Primary Hyperlipidemia
(Mean* % Change from Untreated Baseline†)

<table>
<thead>
<tr>
<th>Treatment</th>
<th>(Daily Dose)</th>
<th>N</th>
<th>Total-C</th>
<th>LDL-C</th>
<th>Apo B</th>
<th>HDL-C</th>
<th>TG*</th>
<th>Non-HDL-C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pooled data (All VYTORIN doses)‡</td>
<td></td>
<td>609</td>
<td>-38</td>
<td>-53</td>
<td>-42</td>
<td>+7</td>
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<td>-49</td>
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<td>Pooled data (All simvastatin doses)‡</td>
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<td>622</td>
<td>-28</td>
<td>-39</td>
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<td>-36</td>
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<td>-18</td>
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<tr>
<td>Placebo</td>
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<td>-2</td>
<td>0</td>
<td>0</td>
<td>-2</td>
<td>-2</td>
</tr>
<tr>
<td>VYTORIN by dose</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10/10</td>
<td></td>
<td>152</td>
<td>-31</td>
<td>-45</td>
<td>-35</td>
<td>+8</td>
<td>-23</td>
<td>-41</td>
</tr>
<tr>
<td>10/20</td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>10 mg</td>
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<td>158</td>
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<td>+5</td>
<td>-17</td>
<td>-30</td>
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<tr>
<td>20 mg</td>
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<td>40 mg</td>
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<td>-21</td>
<td>-38</td>
</tr>
<tr>
<td>80 mg</td>
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<td>158</td>
<td>-35</td>
<td>-49</td>
<td>-39</td>
<td>+7</td>
<td>-27</td>
<td>-45</td>
</tr>
</tbody>
</table>

* For triglycerides, median % change from baseline.
† Baseline - on no lipid-lowering drug.
‡ VYTORIN doses pooled (10/10-10/80) significantly reduced total-C, LDL-C, Apo B, TG, and non-HDL-C compared to simvastatin and significantly increased HDL-C compared to placebo.

In a multicenter, double-blind, controlled, 23-week study, 710 patients with known CHD or CHD risk equivalents, as defined by the NCEP ATP III guidelines, and an LDL-C ≥130 mg/dL were randomized to one of four treatment groups: coadministered ezetimibe and simvastatin equivalent to VYTORIN (10/10, 10/20, and 10/40) or simvastatin 20 mg. Patients not reaching an LDL-C <100 mg/dL had their simvastatin dose titrated at 6-week intervals to a maximal dose of 80 mg.

At Week 5, the LDL-C reductions with VYTORIN 10/10, 10/20, or 10/40 were significantly larger than with simvastatin 20 mg (see Table 8).

Table 8: Response to VYTORIN after 5 Weeks in Patients with CHD or CHD Risk Equivalents and an LDL-C ≥130 mg/dL

<table>
<thead>
<tr>
<th></th>
<th>Simvastatin 20 mg</th>
<th>VYTORIN 10/10</th>
<th>VYTORIN 10/20</th>
<th>VYTORIN 10/40</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>253</td>
<td>251</td>
<td>109</td>
<td>97</td>
</tr>
<tr>
<td>Mean baseline LDL-C</td>
<td>174</td>
<td>165</td>
<td>167</td>
<td>171</td>
</tr>
<tr>
<td>Percent change LDL-C</td>
<td>-38</td>
<td>-47</td>
<td>-53</td>
<td>-59</td>
</tr>
</tbody>
</table>

In a multicenter, double-blind, 6-week study, 1902 patients with primary hyperlipidemia, who had not met their NCEP ATP III target LDL-C goal, were randomized to one of eight treatment groups: VYTORIN (10/10, 10/20, 10/40, or 10/80) or atorvastatin (10 mg, 20 mg, 40 mg, or 80 mg).

Across the dosage range, when patients receiving VYTORIN were compared to those receiving milligram-equivalent statin doses of atorvastatin, VYTORIN lowered total-C, LDL-C, Apo B, and non-HDL-C significantly more than atorvastatin. Only the 10/40 mg and 10/80 mg VYTORIN doses increased HDL-C significantly more than the corresponding milligram-equivalent statin dose of atorvastatin. The effects of VYTORIN on TG were similar to the effects seen with atorvastatin. (See Table 9.)
Table 9: Response to VYTORIN and Atorvastatin in Patients with Primary Hyperlipidemia
(Mean* % Change from Untreated Baseline†)

<table>
<thead>
<tr>
<th>Treatment</th>
<th>(Daily Dose)</th>
<th>N</th>
<th>Total-C‡</th>
<th>LDL-C‡</th>
<th>Apo B‡</th>
<th>HDL-C</th>
<th>TG*</th>
<th>Non-HDL-C‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>VYTORIN by dose</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10/10</td>
<td>230</td>
<td>-34§</td>
<td>-47§</td>
<td>-37§</td>
<td>+8</td>
<td>-26</td>
<td>-43§</td>
<td></td>
</tr>
<tr>
<td>10/20</td>
<td>233</td>
<td>-37§</td>
<td>-51§</td>
<td>-40§</td>
<td>+7</td>
<td>-25</td>
<td>-46§</td>
<td></td>
</tr>
<tr>
<td>10/40</td>
<td>236</td>
<td>-41§</td>
<td>-57§</td>
<td>-46§</td>
<td>+9§</td>
<td>-27</td>
<td>-52§</td>
<td></td>
</tr>
<tr>
<td>10/80</td>
<td>224</td>
<td>-43§</td>
<td>-59§</td>
<td>-48§</td>
<td>+8§</td>
<td>-31</td>
<td>-54§</td>
<td></td>
</tr>
<tr>
<td>Atorvastatin by dose</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10 mg</td>
<td>235</td>
<td>-27</td>
<td>-36</td>
<td>-31</td>
<td>+7</td>
<td>-21</td>
<td>-34</td>
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<tr>
<td>20 mg</td>
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<td>-32</td>
<td>-44</td>
<td>-37</td>
<td>+5</td>
<td>-25</td>
<td>-41</td>
<td></td>
</tr>
<tr>
<td>40 mg</td>
<td>232</td>
<td>-36</td>
<td>-48</td>
<td>-40</td>
<td>+4</td>
<td>-24</td>
<td>-45</td>
<td></td>
</tr>
<tr>
<td>80 mg</td>
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<td>-53</td>
<td>-44</td>
<td>+1</td>
<td>-32</td>
<td>-50</td>
<td></td>
</tr>
</tbody>
</table>

* For triglycerides, median % change from baseline.
† Baseline - on no lipid-lowering drug.
‡ VYTORIN doses pooled (10/10-10/80) provided significantly greater reductions in total-C, LDL-C, Apo B, and non-HDL-C compared to atorvastatin doses pooled (10-80).
§ p<0.05 for difference with atorvastatin at equal mg doses of the simvastatin component.

In a multicenter, double-blind, 24-week, forced-titration study, 788 patients with primary hyperlipidemia, who had not met their NCEP ATP III target LDL-C goal, were randomized to receive coadministered ezetimibe and simvastatin equivalent to VYTORIN (10/10 and 10/20) or atorvastatin 10 mg. For all three treatment groups, the dose of the statin was titrated at 6-week intervals to 80 mg. At each pre-specified dose comparison, VYTORIN lowered LDL-C to a greater degree than atorvastatin (see Table 10).
Table 10: Response to VYTORIN and Atorvastatin in Patients with Primary Hyperlipidemia
(Mean \* % Change from Untreated Baseline\(^\ddagger\))

<table>
<thead>
<tr>
<th>Treatment</th>
<th>N</th>
<th>Total-C</th>
<th>LDL-C</th>
<th>Apo B</th>
<th>HDL-C</th>
<th>TG*</th>
<th>Non-HDL-C</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Week 6</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Atorvastatin 10 mg(^\ddagger)</td>
<td>262</td>
<td>-28</td>
<td>-37</td>
<td>-32</td>
<td>5</td>
<td>-23</td>
<td>-35</td>
</tr>
<tr>
<td>VYTORIN 10/10(^\ddagger)</td>
<td>263</td>
<td>-34(^\ddagger)</td>
<td>-46(^\ddagger)</td>
<td>-38(^\ddagger)</td>
<td>8(^\ddagger)</td>
<td>-26</td>
<td>-43(^\ddagger)</td>
</tr>
<tr>
<td>VYTORIN 10/20(^\ddagger)</td>
<td>263</td>
<td>-36(^\ddagger)</td>
<td>-50(^\ddagger)</td>
<td>-41(^\ddagger)</td>
<td>-25</td>
<td>-26</td>
<td>-46(^\ddagger)</td>
</tr>
<tr>
<td><strong>Week 12</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Atorvastatin 20 mg</td>
<td>246</td>
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<td>-44</td>
<td>-38</td>
<td>7</td>
<td>-28</td>
<td>-42</td>
</tr>
<tr>
<td>VYTORIN 10/20</td>
<td>250</td>
<td>-37(^\ddagger)</td>
<td>-50(^\ddagger)</td>
<td>-41(^\ddagger)</td>
<td>-28</td>
<td>-46(^\ddagger)</td>
<td></td>
</tr>
<tr>
<td>VYTORIN 10/40</td>
<td>252</td>
<td>-39(^\ddagger)</td>
<td>-54(^\ddagger)</td>
<td>-45(^\ddagger)</td>
<td>12(^\ddagger)</td>
<td>-50</td>
<td></td>
</tr>
<tr>
<td><strong>Week 18</strong></td>
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<td></td>
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<td></td>
</tr>
<tr>
<td>Atorvastatin 40 mg</td>
<td>237</td>
<td>-37</td>
<td>-49</td>
<td>-42</td>
<td>8</td>
<td>-31</td>
<td>-47</td>
</tr>
<tr>
<td>VYTORIN 10/40(^\ddagger)</td>
<td>482</td>
<td>-40(^\ddagger)</td>
<td>-56(^\ddagger)</td>
<td>-45(^\ddagger)</td>
<td>11(^\ddagger)</td>
<td>-32</td>
<td>-52(^\ddagger)</td>
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<tr>
<td><strong>Week 24</strong></td>
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<tr>
<td>Atorvastatin 80 mg</td>
<td>228</td>
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<td>-45</td>
<td>6</td>
<td>-35</td>
<td>-50</td>
</tr>
<tr>
<td>VYTORIN 10/80(^\ddagger)</td>
<td>459</td>
<td>-43(^\ddagger)</td>
<td>-59(^\ddagger)</td>
<td>-49(^\ddagger)</td>
<td>12(^\ddagger)</td>
<td>-35</td>
<td>-55(^\ddagger)</td>
</tr>
</tbody>
</table>

\* For triglycerides, median % change from baseline.
\(^\ddagger\) Baseline - on no lipid-lowering drug.
\(^\ddagger\) Atorvastatin: 10 mg start dose titrated to 20 mg, 40 mg, and 80 mg through Weeks 6, 12, 18, and 24.
\(^\ddagger\) VYTORIN: 10/10 start dose titrated to 10/20, 10/40, and 10/80 through Weeks 6, 12, 18, and 24.
\(\ddagger\) p<0.05 for difference with atorvastatin in the specified week.
\(^\ddagger\) VYTORIN: 10/20 start dose titrated to 10/40, 10/40, and 10/80 through Weeks 6, 12, 18, and 24.
\(^\ddagger\) Data pooled for common doses of VYTORIN at Weeks 18 and 24.

In a multicenter, double-blind, 6-week study, 2959 patients with primary hyperlipidemia, who had not met their NCEP ATP III target LDL-C goal, were randomized to one of six treatment groups: VYTORIN (10/20, 10/40, or 10/80) or rosuvastatin (10 mg, 20 mg, or 40 mg).

The effects of VYTORIN and rosuvastatin on total-C, LDL-C, Apo B, TG, non-HDL-C and HDL-C are shown in Table 11.
Table 11: Response to VYTORIN and Rosuvastatin in Patients with Primary Hyperlipidemia
(Mean* % Change from Untreated Baseline†)

<table>
<thead>
<tr>
<th>Treatment (Daily Dose)</th>
<th>N</th>
<th>Total-C‡</th>
<th>LDL-C‡</th>
<th>Apo B‡</th>
<th>HDL-C</th>
<th>TG*</th>
<th>Non-HDL-C‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>VYTORIN by dose</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10/20</td>
<td>476</td>
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<td>-47§</td>
</tr>
<tr>
<td>10/40</td>
<td>477</td>
<td>-39¶</td>
<td>-55¶</td>
<td>-44¶</td>
<td>+8</td>
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<tr>
<td>10/80</td>
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<td>-50#</td>
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<td>-56#</td>
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<td>Rosuvastatin by dose</td>
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<td>-37</td>
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<td>-42</td>
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<tr>
<td>20 mg</td>
<td>478</td>
<td>-37</td>
<td>-52</td>
<td>-43</td>
<td>+8</td>
<td>-26</td>
<td>-48</td>
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<tr>
<td>40 mg</td>
<td>475</td>
<td>-41</td>
<td>-57</td>
<td>-47</td>
<td>+8</td>
<td>-28</td>
<td>-52</td>
</tr>
</tbody>
</table>

* For triglycerides, median % change from baseline.
† Baseline - on no lipid-lowering drug.
‡ VYTORIN doses pooled (10/20-10/80) provided significantly greater reductions in total-C, LDL-C, Apo B, and non-HDL-C compared to rosuvastatin doses pooled (10-40 mg).
§ p<0.05 vs. rosuvastatin 10 mg.
¶ p<0.05 vs. rosuvastatin 20 mg.
# p<0.05 vs. rosuvastatin 40 mg.

In a multicenter, double-blind, 24-week trial, 214 patients with type 2 diabetes mellitus treated with thiazolidinediones (rosiglitazone or pioglitazone) for a minimum of 3 months and simvastatin 20 mg for a minimum of 6 weeks were randomized to receive either simvastatin 40 mg or the coadministered active ingredients equivalent to VYTORIN 10/20. The median LDL-C and HbA1c levels at baseline were 89 mg/dL and 7.1%, respectively.

VYTORIN 10/20 was significantly more effective than doubling the dose of simvastatin to 40 mg. The median percent changes from baseline for VYTORIN vs. simvastatin were: LDL-C -25% and -5%; total-C -16% and -5%; Apo B -19% and -5%; and non-HDL-C -23% and -5%. Results for HDL-C and TG between the two treatment groups were not significantly different.

**Ezetimibe**

In two multicenter, double-blind, placebo-controlled, 12-week studies in 1719 patients with primary hyperlipidemia, ezetimibe significantly lowered total-C (-13%), LDL-C (-19%), Apo B (-14%), and TG (-8%), and increased HDL-C (+3%) compared to placebo. Reduction in LDL-C was consistent across age, sex, and baseline LDL-C.

**Simvastatin**

In two large, placebo-controlled clinical trials, the Scandinavian Simvastatin Survival Study (N=4,444 patients) and the Heart Protection Study (N=20,536 patients), the effects of treatment with simvastatin were assessed in patients at high risk of coronary events because of existing coronary heart disease, diabetes, peripheral vessel disease, history of stroke or other cerebrovascular disease. Simvastatin was proven to reduce: the risk of total mortality by reducing CHD deaths; the risk of non-fatal myocardial infarction and stroke; and the need for coronary and non-coronary revascularization procedures.

No incremental benefit of VYTORIN on cardiovascular morbidity and mortality over and above that demonstrated for simvastatin has been established.

**14.2 Homozygous Familial Hypercholesterolemia (HoFH)**

A double-blind, randomized, 12-week study was performed in patients with a clinical and/or genotypic diagnosis of HoFH. Data were analyzed from a subgroup of patients (n=14) receiving simvastatin 40 mg at baseline. Increasing the dose of simvastatin from 40 to 80 mg (n=5) produced a reduction of LDL-C of 13% from baseline on simvastatin 40 mg. Coadministered ezetimibe and simvastatin equivalent to VYTORIN (10/40 and 10/80 pooled, n=9), produced a reduction of LDL-C of 23% from baseline on simvastatin 40 mg. In those patients coadministered ezetimibe and simvastatin equivalent to VYTORIN (10/80, n=5), a reduction of LDL-C of 29% from baseline on simvastatin 40 mg was produced.
14.3 Chronic Kidney Disease (CKD)

The Study of Heart and Renal Protection (SHARP) was a multinational, randomized, placebo-controlled, double-blind trial that investigated the effect of VYTORIN on the time to a first major vascular event (MVE) among 9438 patients with moderate to severe chronic kidney disease (approximately one-third on dialysis at baseline) who did not have a history of myocardial infarction or coronary revascularization. An MVE was defined as nonfatal MI, cardiac death, stroke, or any revascularization procedure. Patients were allocated to treatment using a method that took into account the distribution of 8 important baseline characteristics of patients already enrolled and minimized the imbalance of those characteristics across the groups.

For the first year, 9438 patients were allocated 4:4:1, to VYTORIN 10/20, placebo, or simvastatin 20 mg daily, respectively. The 1-year simvastatin arm enabled the comparison of VYTORIN to simvastatin with regard to safety and effect on lipid levels. At 1 year the simvastatin-only arm was re-allocated 1:1 to VYTORIN 10/20 or placebo. A total of 9270 patients were ever allocated to VYTORIN 10/20 (n=4650) or placebo (n=4620) during the trial. The median follow-up duration was 4.9 years. Patients had a mean age of 61 years; 63% were male, 72% were Caucasian, and 23% were diabetic; and, for those not on dialysis at baseline, the median serum creatinine was 2.5 mg/dL and the median estimated glomerular filtration rate (eGFR) was 25.6 mL/min/1.73 m², with 94% of patients having an eGFR < 45 mL/min/1.73m². Eligibility did not depend on lipid levels. Mean LDL-C at baseline was 108 mg/dL. At 1 year, the mean LDL-C was 26% lower in the simvastatin arm and 38% lower in the VYTORIN arm relative to placebo. At the midpoint of the study (2.5 years), the mean LDL-C was 32% lower for VYTORIN relative to placebo. Patients no longer taking study medication were included in all lipid measurements.

In the primary intent-to-treat analysis, 639 (15.2%) of 4193 patients initially allocated to VYTORIN and 749 (17.9%) of 4191 patients initially allocated to placebo experienced an MVE. This corresponded to a relative risk reduction of 16% (p=0.001) (see Figure 1). Similarly, 526 (11.3%) of 4650 patients ever allocated to VYTORIN and 619 (13.4%) of 4620 patients ever allocated to placebo experienced a major atherosclerotic event (MAE; a subset of the MVE composite that excluded non-coronary cardiac deaths and hemorrhagic stroke), corresponding to a relative risk reduction of 17% (p=0.002). The trial demonstrated that treatment with VYTORIN 10/20 mg versus placebo reduced the risk for MVE and MAE in this CKD population. The study design precluded drawing conclusions regarding the independent contribution of either ezetimibe or simvastatin to the observed effect.

The treatment effect of VYTORIN on MVE was attenuated among patients on dialysis at baseline compared with those not on dialysis at baseline. Among 3023 patients on dialysis at baseline, VYTORIN reduced the risk of MVE by 6% (RR 0.94: 95% CI 0.80-1.09) compared with 22% (RR 0.78: 95% CI 0.69-0.89) among 6247 patients not on dialysis at baseline (interaction P=0.08).

Figure 1: Effect of VYTORIN on the Primary Endpoint of Risk of Major Vascular Events
The individual components of MVE in all patients ever allocated to VYTORIN or placebo are presented in Table 12.

Table 12: Number of First Events for Each Component of the Major Vascular Event Composite Endpoint in SHARP*

<table>
<thead>
<tr>
<th>Outcome</th>
<th>VYTORIN 10/20 (N=4650)</th>
<th>Placebo (N=4620)</th>
<th>Risk Ratio (95% CI)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Major Vascular Events</td>
<td>701 (15.1%)</td>
<td>814 (17.6%)</td>
<td>0.85 (0.77-0.94)</td>
<td>0.001</td>
</tr>
<tr>
<td>Nonfatal MI</td>
<td>134 (2.9%)</td>
<td>159 (3.4%)</td>
<td>0.84 (0.66-1.05)</td>
<td>0.12</td>
</tr>
<tr>
<td>Cardiac Death</td>
<td>253 (5.4%)</td>
<td>272 (5.9%)</td>
<td>0.93 (0.78-1.10)</td>
<td>0.38</td>
</tr>
<tr>
<td>Any Stroke</td>
<td>171 (3.7%)</td>
<td>210 (4.5%)</td>
<td>0.81 (0.66-0.99)</td>
<td>0.038</td>
</tr>
<tr>
<td>Non-hemorrhagic Stroke</td>
<td>131 (2.8%)</td>
<td>174 (3.8%)</td>
<td>0.75 (0.60-0.94)</td>
<td>0.011</td>
</tr>
<tr>
<td>Hemorrhagic Stroke</td>
<td>45 (1.0%)</td>
<td>37 (0.8%)</td>
<td>1.21 (0.78-1.86)</td>
<td>0.40</td>
</tr>
<tr>
<td>Any Revascularization</td>
<td>284 (6.1%)</td>
<td>352 (7.6%)</td>
<td>0.79 (0.68-0.93)</td>
<td>0.004</td>
</tr>
</tbody>
</table>

*Intention-to-treat analysis on all SHARP patients ever allocated to VYTORIN or placebo.

Among patients not on dialysis at baseline, VYTORIN did not reduce the risk of progressing to end-stage renal disease compared with placebo (RR 0.97: 95% CI 0.89-1.05).

16 HOW SUPPLIED/STORAGE AND HANDLING

No. 3873 — Tablets VYTORIN 10/10 are white to off-white capsule-shaped tablets with code “311” on one side.

They are supplied as follows:

NDC 66582-311-31 bottles of 30
NDC 66582-311-54 bottles of 90
NDC 66582-311-87 bottles of 10,000 (If repackaged in blisters, then opaque or light-resistant blisters should be used.)

No. 3874 — Tablets VYTORIN 10/20 are white to off-white capsule-shaped tablets with code “312” on one side.

They are supplied as follows:

NDC 66582-312-31 bottles of 30
NDC 66582-312-54 bottles of 90

No. 3875 — Tablets VYTORIN 10/40 are white to off-white capsule-shaped tablets with code “313” on one side.

They are supplied as follows:

NDC 66582-313-31 bottles of 30
NDC 66582-313-54 bottles of 90

No. 3876 — Tablets VYTORIN 10/80 are white to off-white capsule-shaped tablets with code “315” on one side.

They are supplied as follows:

NDC 66582-315-31 bottles of 30
NDC 66582-315-54 bottles of 90

**Storage**

Store at 20-25°C (68-77°F). [See USP Controlled Room Temperature.] Keep container tightly closed.

**Storage of 10,000, 5000, and 2500 count bottles**

Store bottle of 10,000 VYTORIN 10/10 and 10/20, 5000 VYTORIN 10/40, and 2500 VYTORIN 10/80 capsule-shaped tablets at 20-25°C (68-77°F). [See USP Controlled Room Temperature.] Store in original container until time of use. When product container is subdivided, repackage into a tightly-closed, light-resistant container. Entire contents must be repackaged immediately upon opening.
17 PATIENT COUNSELING INFORMATION

Advise the patient to read the FDA-approved patient labeling (Patient Information).

Patients should be advised to adhere to their National Cholesterol Education Program (NCEP)-recommended diet, a regular exercise program, and periodic testing of a fasting lipid panel.

Patients should be advised about substances they should not take concomitantly with VYTORIN [see Contraindications (4) and Warnings and Precautions (5.1)]. Patients should also be advised to inform other healthcare professionals prescribing a new medication or increasing the dose of an existing medication that they are taking VYTORIN.

17.1 Muscle Pain

All patients starting therapy with VYTORIN should be advised of the risk of myopathy, including rhabdomyolysis, and told to report promptly any unexplained muscle pain, tenderness or weakness particularly if accompanied by malaise or fever or if these muscle signs or symptoms persist after discontinuing VYTORIN. Patients using the 10/80-mg dose should be informed that the risk of myopathy, including rhabdomyolysis, is increased with the use of the 10/80-mg dose. The risk of myopathy, including rhabdomyolysis, occurring with use of VYTORIN is increased when taking certain types of medication or consuming grapefruit juice. Patients should discuss all medication, both prescription and over the counter, with their healthcare professional.

17.2 Liver Enzymes

It is recommended that liver function tests be performed before the initiation of VYTORIN, and thereafter when clinically indicated. All patients treated with VYTORIN should be advised to report promptly any symptoms that may indicate liver injury, including fatigue, anorexia, right upper abdominal discomfort, dark urine or jaundice.

17.3 Pregnancy

Women of childbearing age should be advised to use an effective method of birth control to prevent pregnancy while using VYTORIN. Discuss future pregnancy plans with your patients, and discuss when to stop taking VYTORIN if they are trying to conceive. Patients should be advised that if they become pregnant they should stop taking VYTORIN and call their healthcare professional.

17.4 Breastfeeding

Women who are breastfeeding should be advised to not use VYTORIN. Patients who have a lipid disorder and are breastfeeding should be advised to discuss the options with their healthcare professional.

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