(2.5.7.2)
Wafarii: Obtain INR prior to starting rosuvastatin tablets. Monitor INR frequently until stable upon initiation, dose titration or discontinuation. (7.3)
INSF IM SPECIFIC POPULATIONS USE IN SPECIFIC POPULATIONS ----

•	17 for DATIENT COUNCELING INCOMMATION and EDA approved national labeling
•	Lactation: Breastfeeding not recommended during treatment with rosuvastatin tablets.

Consider if the benefit of using fibrates concomitantly with resunant outweights the increased risk of myopathy and shabdomyolysis. If co-decided, monitor patients for signs and symptoms of myopathy, par initiation of therapy and during upward dose thration of either drug.

microarrange (and Dist). It is used reproduction statistics, she adverse developmental ethics, were observed in progrant statis or rabbits to insert reproduction statistics, she adverse development in the statistic production of the branch on programs of the microarrange statistic production of the branch on programs of the statistic production of the district productio

keimal Data 1 demale rats given 5, 15 and 50 mg/kg/day before mating and continuing through to gestation day 7 resulted n decinased fetal body weight (fernale pups) and delayed ossification at 50 mg/kg/day (10 times the human

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ULL	PRESCRIBING	INFORMATION: C	ONTENTS*
1	INDICATIONS	AND USAGE	

- DISSACE AND ADMINISTRATION

  2. Recommended Dosage and Administration Information

  2. Recommended Dosage and Administration

  3. Recommended Dosage in Pedatire Pleatents

  4. Dosing in Asian Patients

  4. Dosing in Asian Patients

  5. Recommended Dosage in Patients with Renal Impairment

  5. Dosage and Administration Modifications Due to Drug Inter

  DOSAGE FORMS AND STRENTIS

  CONTRAINDICATIONS

  CONTRAINDICATIONS

- CONTRAINDICATIONS

  5.1 Myopathy and Rhabdomyolysis

  1. mnane-Mediated Necrotizing Myopathy
  5.3 Hepatic Dysfunction
  5.4 Proteinuris and Hematuria
  5.5 Increases in HbA1c and Fasting Serum Gl
- 5.5 Increases in HbA1c and Fast ADVERSE REACTIONS 6.1 Clinical Trials Experience 6.2 Postmarketing Experience

PATHERS with Settler received impairment in the control of the con

- USE IN SPECIFIC POPULATIONS
  8.1 Pregnancy
  8.2 Lactation
  8.4 Pediatric Use
  8.5 Geraturic Use
  8.6 Real Impairment
  8.7 Hepate Impairment
  8.7 Hepate Impairment
  8.8 Anna Patients
  00\*\*CRINICAL PHARMACOLOGY
  12.1 Mechanism of Action
  00\*\*CRINICAL PHARMACOLOGY
  12.1 Pharmacological Pharmacologica

- 12.5 Plasmacogenomics
  NONCLINICAL TOXICOLOGY
  13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility
  CLINICAL STUDIES
  NOW SUPPLIED/STORAGE AND HANDLING
  PATIENT COUNSELING INFORMATION
  \*\*Sections or subsections omitted from the full prescribing info

ILL PRESCRIBING INFORMATION INDICATIONS AND USAGE	Table 2: Advers Placebs-Contro	se Reactions Re olled Trials	ported in >2% o	of Patients Treat	ted with Rosuva	statin Table
soverstatin tablets is indicated:  To reduce the risk of stroke, myccardial interction, and arterial revascularization procedures in adults without established corenary heart disease who are all increased risk of cardiovascular (OV) disease based on age, IsoCRP >2 mg/L, and at least one additional OV risk factor.	Adverse Reactions	Placebo N=382 %	Rosuvastašn 5 mg N=291 %	Resuvastatin 10 mg N=283 %	Rosuvastatin 20 mg N=64 %	Resuvast 40 mg N=106 %
As an adjunct to diet to:	Headache	5.0	5.5	4.9	3.1	8.5

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- conformation events.

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	Modifications Due to Drug Interactions
Concomitantly Used Drug	Resuvastatin tablets Dosage Medifications
Cyclosporine	Do not exceed 5 mg once daily.
Teriflunomide	Do not exceed 10 mg once daily.
Enasidenib	Do not exceed 10 mg once daily.
Capmatinib	Do not exceed 10 mg once daily.
Fostamatinib	Do not exceed 20 mg once daily.
Febuxostat	Do not exceed 20 mg once daily.
Gernfibrozii	Avoid concomitant use. If used concomitantly, initiate at 5 mg once daily and do not exceed 10 mg once daily.
Tafamidis	Avoid concomitant use. If used concomitantly, initiate at 5 mg once daily and do not exceed 20 mg once daily.
Antiviral Medications	
o Sofbuvir/velpatasvir/voxilaprevir o Ledipasvir/sofosbuvir	Concomitant use not recommended.
o Simeprevir o Dasabuviciombitasvici paritapreviritionavir o Elbasvici/Grazoprevir o Sofosbuvic/Velpatasvir	Initiate at 5 mg once daily. Do not exceed 10 mg once daily

 Darnistriide
 Darnistriide
 Darnistriide
 Darnistriide
 Regoralenib
 Regoralenib Resuscitatin Tablets Administration Medifications Due to Drup interactions
When taking resussatatin tablets with an aluminum and magnesium hydroxide combin
resussatatin tablets at least 2 hours before the antacid (see Ang Interactions (7.2)).

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Rosuvastatili tablets may cause myopathy (muscle pain, tenderness, or weakness associated with elevated creatine kinase (DK)) and materiomyolysis. Acute kidnby (painy secondary to myoplobinuria and rare fatalities have occurred as a result of midelomyolysis with statiss; including monoacration fatalities.

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with unoplained persistent proteinaria and/or hematuria during metine urinalysis testing.

5.5. Increases in HANL and Francia Searum Gloscote Levels
Increases in HANL and Francia Searum Gloscote levels have been reported with statins, including rosuvastatio
tablets. Based on clinical third data with rosuvastation bablets, in some instances these compacts may over
thresholds for the diagnosis of clinicals methods. [Or degree Enrollection 12.1]. Optimize History in missures,

under widely varying conditions, adverse reaction rates observed in the titly compared to rates in the clinical trials of another drug and may not utilize. of a drug cannot be orrectly compared to name or an administration of so observed in clinical practice. icos reported in 22% of patients in placebo-controlled clinical studies and nown in Table 2. These studies had a treatment duration of up to 12 weeks.

Adverse Reactions	Placebo N=382 %	Rosuvastatin 5 mg N=291 %	Resuvestation 10 mg N=283 %	Rosuvastatin 20 mg N=64 %	Rosuvastatin 40 mg N=106 %	Total Rosuvastatin 5 mg to 40 mg N=744 %
Headache	5.0	5.5	4.9	3.1	8.5	5.5
Nausea	3.1	3.8	3.5	6.3	0	3.4
Myalgia	1.3	3.1	2.1	6.3	1.9	2.8
Asthenia	2.6	2.4	3.2	4.7	0.9	2.7
Constinution	2.4	2.1	2.1	4.7	2.8	2.4

	hypercholesterolemia (HoFH).	Constipation	2.4	2.1	2.1	4.7	2.8	2.4
	As an adjunct to diet for the treatment of adults with:  o Primary dystetalipoproteinemia. o Hypertriggleenidemia.	rash, pruritus, been recorded:	urticaria, and an	ngioedema) and	pancreatitis. The	e following lab	ness, hypersensi oratory abnorma wated creatine i	dities have also shosobokinase
	DOSAGE AND ADMINISTRATION	transaminases, abnormalities	glucose, glutar	myl transpeptid	ase, alkaline ph	osphatase, and	bilirubin; and t	hyrold function
	General Decage and Africational Internation Administer recognition that has a single dose at any time of day, with or without food. The tablet should be suchtioned whose a single dose at any time of day, with or without food. The tablet should be suchtioned whose and the single second of the second second of the Assess LIU.C.—Very me clinically appropriate, as early as 4 weeks after initiating resuscetatin tablets and adjust the closure in recognition of the amount of the second s	In the METEOR mean treatmen placebo are sho	t duration of 1.7 wn in Table 3. se Reactions Re	years. Adverse	reactions report	ed in 22% of pa	-700) or placebo dients and at a r statin Tablets a	ate greater than
2	Recommended Dosage in Adult Patients The dosage range for resunstatin tablets is 5 to 40 mg orally once daily.	Adver	se Reactions		Placebo N=281 %		Rosuvastatin tal N=700	
	The recommended dose of resulastatin tablets depends on a patient's indication for usage, LDL-C, and individual risk for cardiovascular events.	Myalgia			12.1		12.7	
3	Recommended Dosage in Pediatric Patients	Arthralgia			7.1		10.1	
	s in Pertiatric Patients 8 Years of Age and Older with HeFH commended dosage range is 5 mg to 10 mg onlike once daily in patients aged 8 years to less than 10	Headache			5.3		6.4	
275 Z	commitment detailer range is 5 mg to 10 mg otary once tarry in patients aged 6 years to less than 10 md 5 mg to 20 mg orally once daily in patients aged 10 years and older.  In Rediatric Patients 7 Years of Ace and Older with HoFH	Dizziness			2.8		4.0	
	commended dosage is 20 mg grally once daily.	Increased 0	PK		0.7		2.6	
	Dosing in Asian Patients	Abdominal	pain		1.8		2.4	
risi	rosuvastatin tablets at 5 mg once daily due to increased rosuvastatin plasma concentrations. Consider ks and benefits of rosuvastatin tablets when treating Asian patients not adequately controlled at doses up	ALT greate	rthan 3x ULN		0.7		2.2	
20	mg once daily [see Warnings and Procautions (S.1), Use in Specific Populations (R.R), and Clinical	Frequency reco	orded as abnorm	al laboratory val	ue.			

tablets-treated versus placebo-treater Adverse reactions reported in ×2% or	1 patients <i>(see Clinical Studies (14</i> )). 1 patients and at a rate greater than pl	s significantly higher in resuvestant acebo are shown in Table 4. nsuvastatin Tablets and > Placebo in	needs of the individual patient.
Adverse Reactions	Placebo N=8901 %	Rosuvastatin tablets 20 mg N=8901 %	Resuvastatin tablets decreases synthesis of chelesterol and possibly other biologically active substances derived from chalesterol; therefore, resuvastatin tablets may cause fetal harm when administered to pregnant patients based on the mechanism of action (see <u>Clinical Pharmacology 1/2 1/1</u> ), in addition, treatment of
Myalgia	6.6	7.6	hyperlipidemia is not generally necessary during pregnancy. Atherosciensis is a chronic process and the discontinuation of lipid-lowering drups during pregnancy should have little impact on the outcome of long-term
Arthralgia	3.2	3.8	therapy of primary hyperlipidemia for most patients.
Constipation	3.0	3.3	Available data from case series and prospective and retrospective observational cohort studies over decades of use with statins in pregnant women have not identified a drug-associated risk of major congenital
Diabetes mellitus	2.3	2.8	malformations. Published data from prospective and retrospective observational cohort studies with resuvastatin tablets use in pregnant women are insufficient to determine if there is a drug-associated risk of
Nausea	2.3	2.4	miscarriage (see Data). In animal reproduction studies, no adverse developmental effects were observed in pregnant rats or rabbits
Padiatric Patients with HoFH			orally administered resuvastatin during the period of organogenesis at doses that resulted in systemic

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Hepotobilisy Disorders: artiralgia, rare reports of immune-mediated necro statil use

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Cyclosponne	
Clinical Impact:	Cyclosporine increased resuvastatin exposure 7-fold. The risk of myopathy and rhabdomyolysis is increased with concomitant use of cyclosporine or gentifibrazil with resuvastatin tablets.
Intervention:	If used concomitantly, do not exceed a dose of resuvastatin tablets 5 mg once daily.
Teriffunomide	
Clinical Impact:	Teriflunomide increased resuvastatin exposure more than 2.5-fold. The risk of myopathy and rhabdomyolysis is increased with concomitant use.
Intervention:	In patients taking teriflunomide, do not exceed a dose of resuvastatin tablets 10 mg once daily.
Enasidenib	
Clinical Impact:	Enasidenib increased rosuvastatin exposure more than 2.4-fold. The risk of myopathy and shabdomyolysis is increased with concomitant use.
Intervention:	In patients taking enasidentb, do not exceed a dose of rosuvastatin tablets 10 mg once daily.
Capmatinib	
Olinical Impact:	Capmatinib increased resuvastatin exposure more than 2.1-fold. The risk of myopathy and rhabdomyolysis is increased with concomitant use.
Intervention:	In patients taking capmatinib, do not exceed a dose of rosuvastatin tablets 10 mg once daily.

Fostmatishi increased rosuvastatin exposure more than 2.0-fold. The risk myopathy and rhabdomyolysis is increased with concomitant use.
 In patients taking fostamatinib, do not exceed a dose of rosuvastatin tablet once daily.

Avoid concentrat use of gerntorcoll with rosinustatin tablets.

Avoid concentrat use of gentificati with resunstatin tablets. If used concentrational institute resuscatatin tablets if my once daily and do not exceed a dose of rosinusatatin tablets 10 mg once daily. Resuvastatin plasma levels were significantly increased with concomitant administration of many anti-viral drugs, which increases the risk of myopathy and

Fine Charage and Administrations (2.4), and Disord Plannamings (12.3).

10 UNERGOSAIE
No specific artificities for researchain tablets are known. Hemodalayist does not signific or researchain Contact Prisess Central (4.60.0222-1222) for latest recommendations.

11 DESCRIPTION
PROMODER OF White One Prechividation Control (4.60.022). inflition and in the many are man series, when the excession are not in regionary and inflition and inflited \_\_\_\_\_ with \_\_\_\_\_ with tablets initiate with reservantatin tablets 5 mg once daily, and do not exceed a doze of reservantatin tablets 10 mg once daily. The chemical name for resolvation cacking 18 at 9 primiting values of 6 + 10 primiting values of 6 primiting v sical frepact: Dareletamide increased resuxestatin exposure more than 5 fold. The risk of myopati and makdomyolysis is increased with concombant use. In patients taking dareletamide, do not exceed a dose of resuxestatin tablets 5 mg once daily. egoraferib

linical Impact: Regoraferib increased rosuvastatin exposure and may increase the risk of myopathy.

newvercer: In patients taking reporaferib, do not exceed a dose of rosuvastatin tablets 10 mg

The empirical formula for resurractatin calcium USP is (C\_H\_RN\_0.5), ca and the molecular weight is 1001.14. Resurractatin calcium USP is a Withe or almost white, bytypicacopic powder that is freely soluble in methylene otheride, slightly soluble in water practically insoluble in arrhydrous ethand. Resurractatin calcium is a hydrochilic comound with a cutificial coefficient increductation of 19.5 and 47.5 is 4.44.47.5 is Afterwarms. coce daily.

Fenestbrakes (e.g., tenestbrake and fenestbric acids)

Afficial insect: Fibrates may cause myopathy when given alone. The risk of myopathy and insect acids and the second and the conventional use of fibrates with recurrents. Researchtie tablets for onal use contain researchtie 5 mg, 10 mg, 20 mg, or 40 mg (equivalent to 5.2 mg, 10 mg, 20 mg, or 40 mg (equivalent to 5.2 mg, 10 mg, 20 mg, or 40 mg (equivalent to 5.2 mg, 10 mg, 20 mg, 2

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Absoration in Indicate Institution of the Indicate Institution of institution or institution were reached 3 to 5 hours belowing self-coloring both Co., and AUC Increased in approximate projection to consisted while side on The coloring or control good periodication. Considerating SNRs has All of increased and self-coloring SRs of sold.

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Americantation of conventable habits with tool did not affect the AUC of resonabilists.

Distribution

Mean volume of distribution at steady-state of rosuvastatin is approximately 134 liters. Rosuvastatin is 86% bound to plasma proteins, mostly albumin. This binding is reversible and independent of plasma concentrations.

Security interest products may deploy this besides the models and trained secretary of memory and construction.

Reconsisting as not extensively conductative and approximately 10% of a implication of construction. The consequence of the cons

Planets with native important in Mild to moderate real impairment (CL, 250 mL/min/1.73 m²) had no influence on plasma concentrations of resuvestatin. However, plasma concentrations of resuvestatin increased to a clinically significant extent (about 3-fals) in plainets with severe real impairment (CL, 260 mL/min/1.73 m²) not receiving hemodialpsis compared with healthy subjects (CL, 260 mL/min/1.73 m²).

with beatily souldest, EL, 500 off, birth 7.73 m<sup>2</sup>). The design of the control o

100% and 21%, respectively, compared with patients with normal liver function.

Drug Internatives Studies
Resulvastatin clearance is not dependent on metabolism by cytochrome P450 3A4 to a clinically significant control.

Researchin is a substrate for contain transporter problem including the hequits uptake transporter organic auto-cracecoping polysection (1871, (MATHER)) and effect transporter bread concer metabone protein (1872, MATHER) and the transporter bread concer metabone protein (1872, MATHER) and the transporter bread concernitation of transportation (1874, MATHER) and the transporter bread concernitations (1874, MATHER) and the transporter bread concernitations (1874, MATHER) and transporter bread concernitations (1874, MATHER) and transporter bread concernitations (1874, MATHER) and transporter (1874, MATHER)

Coadministered drug and dosing regimen		Rosuvastatin	
		Mean Ratio (rate with/withou drug) No Effect-1	
	Dose (mg) <sup>1</sup>	Change in AUC	Change in C
Sotosbuvir/velpatasvir/voxilaprevir (400 mg-100 mg-100 mg) + Voxilaprevir (100 mg) once daily for 15 day	10 mg, single dose	7.39 <sup>2</sup> (6.68-8.18) <sup>2</sup>	18.88° (16.23-21.96)°
Cyclosporine – stable dose required (75 mg – 200 mg BID)	10 mg, QD for 10 days	7.12	117
Darolutamide 600 mg BID, 5 days	5 mg, single dose	5.20	-5'
Regoratenib 160 mg QD, 14 days	5 mg, single dose	3.82	4.8'
Atazanavir/ritonavir combination 300 mg/100 mg QD for 8 days	10 mg	3.1°	72
Simeprevir 150 mg QD, 7 days	10 mg, single dose	2.8° (2.3-3.4)°	3.2° (2.6-3.9)°
Velpatasvir 100 mg once daily	10 mg, single dose	2.69° (2.46-2.94)°	2.61° (2.32-2.92)°
Ombitasvir 25 mg/paritaprevir 150 mg/ ritonavir 100 mg + dasabuvir 400 mg BID	5 mg, single dose	2.59° (2.09·3.21)°	7.13° (5.11-9.96)°
Teriflunomide	Not available	2.51°	2.65
Enasidenib 100 mg QD, 28 days	10 mg, single dose	2.44	3.66
Elbasvir 50 mg/grazoprevir 200 mg once daily	10 mg, single dose	2.26° (1.89-2.69)°	5.49° (4.29-7.04)°
Glecaprevir 400 mg/pibrentasvir 120 mg once daily	5 mg, once daily	2.15° (1.88-2.46)°	5.62° (4.80-6.59)°
Lopinavin'ritonavir combination 400 mg/100 mg BID for 17 days	20 mg, QD for 7 days	2.1 <sup>2</sup> (1.7-2.6) <sup>3</sup>	5° (3.4-6.4)°
Capmatinib 400 mg BID	10 mg, single dose	2.08° (1.56-2.76)°	3.04° (2.36-3.92)°
Fostamatinib 100 mg BID	20 mg, single dose	1.96° (1.77-2.15)°	1.88° (1.69-2.09)°
Febuxostat 120 mg QD for 4 days	10 mg, single dose	1.9° (1.5·2.5)°	2.1° (1.8-2.6)°
Gernfibrozii 600 mg BID for 7 days	80 mg	1.9 <sup>2</sup> (1.6-2.2) <sup>3</sup>	2.1° (1.8/2.7)°
Tafamidis 61 mg BID on Days 1 & 2, followed by QD on Days 3 to 9	10 mg	1.97° (1.68-2.31)	1.86 <sup>2</sup> (1.59-2.16) <sup>3</sup>
Eltrombopag 75 mg QD, 5 days	10 mg	1.6 (1.4-1.7) <sup>2</sup>	2 (1.8-23) <sup>3</sup>
Darunavir 600 mg/ktoravir 100 mg BID, 7 days	10 mg, QD for 7 days	1.5 (1.0·2.1) <sup>3</sup>	2.4 (1.6-3.6) <sup>2</sup>
Tipranavirhitonavir combination 500 mg/200 mg BID for 11 days	10 mg	1.4 (1.2-1.6) <sup>2</sup>	2.2 (1.8-2.7) <sup>3</sup>
Dronedarone 400 mg BID	10 mg	1.4	
traconazole 200 mg QD, 5 days	10 mg or 80 mg	1.4 (1.2-1.6) <sup>2</sup> 1.3 (1.1-1.4) <sup>2</sup>	1.4 (1.2-1.5) <sup>3</sup> 1.2 (0.9-1.4) <sup>3</sup>
Ezetimibe 10 mg 00, 14 days	10 mg, QD for 14 days	12 (0.9-1.6)2	12 (0.8-1.6) <sup>3</sup>
Fosamprenavir/ritonavir 700 mg/100 mg BID for 7 days	10 mg	1.1	1.5
Fenofibrate 67 mg TID for 7 days	10 mg	**	1.2 (1.1+1.3) <sup>2</sup>
Rifampicin 450 mg QD, 7 days	20 mg	**	
Aluminum & magnesium hydroxide combination antazid Administered simultaneously Administered 2 hours apart	40 mg 40 mg	0.5° (0.4-0.5)° 0.8	0.5° (0.4-0.6)° 0.8

(ratio with/without drug) No Effect = 1 Change in AUC R- Warfarin 1.0 (1.0-1.1)<sup>2</sup> S-Warfarin 1.0 (0.9-1.0)<sup>2</sup> S-Wartarin

Oral Contraceptive (ethinyl estradiol 0.035 mg & norgestrel 0.180, 0.215 and 0.250 mg) QD for 21 I

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sny end point was a composite end point consisting of the time-to-first occurrence of any of the major CV events: CV death, nontatal impocadiol infanction, nontatal strole, hospitalization for unstable

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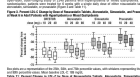


are unablead components of the primary end point are presented in Figure 3. Resovuestatin tables significantly reduced the risk of enotation grounds interrisk no nontrial stroke, and afterful resociationation procedures. There were no significant treatment differences between the resovuestatin tablets and placebo groups for death due to cardiovacular causes or heapplicitations for unstable areains. These was confident bushed extractor common 

	(n-800) (n-800)	(Nacion 20 mg (Nacion) (12097)	Minerca	Probus	Roselvo	1-10%
Cycle (RC)	HOTE	3620040	E30 SLM, GAST	1000		
	27.2.95	W(34)	6300021.124	920	-	
			8363KII,650	1946		
ngra.			ANGLE, CHE	900	-	
orien	77 205	101 (7.0)	EMBRECESS			
֡	organic SICII rate dustr <sup>a</sup> loute at ed regre	Rocks 2 mg   14-80m)   14-80m)   15-80m)   15-80m   15-20m   15-20	In-attent    In-		Nove 2 Feb   Part   Part   Part	Novem 2

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	Treatment Daily Dose					
Treatment	10 mg	20 mg	40 mg	80 mg		
Rosuvestatin	-46'	-521	-55*	_		
Atorvastatin	-37	-43	-48	-51		
Simvastatin	-28	-35	-39	-46		
Prayastatin	-20	-24	-30	_		

Corresponding tastesier erinns are approximately 1.00.

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all patient level in the group treated with reservantatin tablets, 52.1% of patients demonstrated an iscase progression (defined as a negative annualized rate of change), compared to 37.7% of placedo group.

		Rosuvastatin tablets (n=435) LS Mean' (95% CI)	Atomastatin (n=187 LS Mean* (95% CI
Week 6	20 mg	-47% (-49%, -48%)	-38% (-40%, -36%)
Week 12	40 mg	-55% (-57%, -54%)	47% (49%, 45%
Week 18	80 mg	NA .	-52% (-54%, -50%

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Dose (mg)	N	LDL-C	HDL-C	Total-C	Tg'	ApaB
Placebo	46	-1%	+7%	0%	-7%	-2%
5	42	-38%	+4%*	-30%	-13%	-32%
10	44	-45%	+11%	-34%	-15%	-38%
20	44	-50%	+9%"	-39%	16%	-41%

Representative was done charteful in a New-year open-label, reconstruct, distribution-to-good in their desirable of their sound and desirable on their desirable of t in IDL-C from baseline were generally consistent across age groups within the trial as well as experience in both adult and pediatric controlled trials.

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In a super-index Constitution State, Martin patients, (i)—40, 443 years) were evaluated to their responses as in an appreciate Constitution State, Martin State Constitution State Constitu

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Lipid-modifying Etherics of Recoversation Tablets 10 mg and 20 mg in Adult Palients with Primary isoprovincemia (Type III hyperispoproteinemia) After Six Weeks by Median Percent Change (95% Saxeline Min-2).

	(mg/dL)	from baseline (95% Cl) Rosuvastatin tablets 10 mg	
Total-C	342.5	-43.3 (-46.9, = 37.5)	-47.6 (-51.6,-42.8)
Triglycerides	503.5	-40.1 (-44.9, -33.6)	-43.0 (-52.5, -33.1)
Non-HDL-C	294.5	-48.2 (-56.7, -45.6)	-56.4 (-61.4, -48.5)
VLDL-C + IDL-C	209.5	-46.8 (-53.7, -39.4)	-56.2 (-67.7, -43.7)
LDL-C	112.5	-54.4 (-59.1, -47.3)	-57.3 (-59.4, -52.1)
HDL-C	35.5	10.2 (1.9, 12.3)	11.2 (8.3, 20.5)
RLP-C	82.0	-56.4 (-67.1, -49.0)	-64.9 (-74.0, -56.6)
Apo-E	16.0	-42.9 (-46.3, -33.3)	-42.5 (-47.1, -35.6)

In a double-blind, placebo-controlled study in adult patients with baseline TG levels from 273 to 817 mg/dL, cooxvactatin tablets given as a single daily dose (5 to 40 mg) over 6 weeks significantly reduced serum TG levels (Table 16). Table 16: Lipid-Modifying Effect of Resuvastatin Tablets in Adult Patients with Pr After Six Weeks by Median (Min. Max) Percent Change from Buseline to Week 6

Dose	Placebo (n=26)	Resuvastatin tablets 5 mg (n=25)	Rosuvastatin tablets 10 mg (n=23)	Rosavastatin tablets 20 mg (n=27)	Rosuvastatin tablets 40 mg (n=25)
Triglycerides	1 (-40, 72)	-21 (-58, 38)	-37 (-65, 5)	-37 (-72, 11)	-43 (-80, -7)
Non-HDL-C	2 (-13, 19)	-29 (-43, -8)	49 (-59, -20)	-43 (-74, 12)	-51 (-62, -6)
Total-C	1 (-13, 17)	-24 (-40, -4)	-40 (-51, -14)	-34 (-61, -11)	-40 (-51, -4)
LDL-C	5 (-30, 52)	-28 (-71, 2)	-45 (-59, 7)	-31 (-66, 34)	-43 (-61, -3)
HDL-C	-3 (-25, 18)	3 (-38, 33)	8 (-8, 24)	22 (-5, 50)	17 (+14, 63)
6 HOW S	UPPLIED/STORA	AGE AND HANDLIN	6		

Rosuvastati	n Tablets USP are supp		
Strength	How Supplied	NDC Tablet	Description
5 mg	Bottle of 90 tablets Bottle of 500 tablets	NDC 13668-720-90 NDC 13668-720-05	Pink, round, biconvex, beveled edge, film coate tablets debossed with "RS" on one side and pla on other side.
10 mg	Bottle of 90 tablets Bottle of 500 tablets	NDC 13668-721-90 NDC 13668-721-05	Pink, round, biconvex, beveled edge, film coate tablets debossed with "R10" on one side and pi on other side.
20 mg	Bottle of 90 tablets Bottle of 500 tablets	NDC 13668-722-90 NDC 13668-722-05	Pink, round, biconvex, film coated tablets debossed with "R20" on one side and plain on other side.
40 mg	Bottle of 30 tablets	NDC 13668-723-30 MDC 13668-723-05	Pink, oval, biconvex, film-coated tablets deboss

Storage
Store at controlled room temperature, 20°C to 25°C (68°F to 77°F); excursions permitt
30°C (58°F and 86°F) (see USP Controlled Room Temperature). Protect from moistur 30 C (30 P and 60 P) (see con-continued recent interpretation). Patient COUNSELING INFORMATION Advise the patient to read the FDA-approved patient labeling (Patient Information).

ann reconstruct; c-rt, and may reconstruct (z-1).

Highasit Dystanction

Inform patients that resuscitatio tablets may cause liver enzyme elevations and possibly liver failure. Advise
patients to promptly report tabuse, ancreais, right upper abdominal disconflort, dark write or journduce face

Microsine and Processions (S-SE).

Made in India Manufactured for: TORRENT PHARMA INC. Basking Ridge, NJ 07920.

Rev: 12/23, V-03 ROSUVASTATIN (nee soo" va star" in) TABLETS USP allion carefully before you start taking resuvastatin tablets and each time you get assons about reasuvastatin tablets, ask your dector. Only your dector can determin

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