

Alphagan® P 0.1%

(brimonidine tartrate ophthalmic solution) 0.1%

REMEMBER THE GREEN BOTTLE



Use **ALPHAGAN® P 0.1%**
Every Day, the Right Way

INDICATIONS AND USAGE

ALPHAGAN® P (brimonidine tartrate ophthalmic solution) 0.1% or 0.15% is an alpha-adrenergic receptor agonist indicated for the reduction of elevated intraocular pressure (IOP) in patients with open-angle glaucoma or ocular hypertension.

IMPORTANT SAFETY INFORMATION

CONTRAINDICATIONS

Neonates and Infants (under the age of 2 years): ALPHAGAN® P is contraindicated in neonates and infants (under the age of 2 years).

Hypersensitivity Reactions: ALPHAGAN® P is contraindicated in patients who have exhibited a hypersensitivity reaction to any component of this medication in the past.

Please see additional Important Safety Information inside.

A medication diary and helpful tips for using your eyedrops

Remember to use **ALPHAGAN® P 0.1%** every day as directed

Use these tips to help you remember to use your eyedrops:



- Set a daily clock or watch alarm that can remind you to use your eyedrops



- Associate using your eyedrops with other daily routines you've established for yourself, such as brushing your teeth



- Ask a friend or family member to remind you when it's time to use your eyedrops



- Keep this medication diary handy as a reminder to take and track each dose

IMPORTANT SAFETY INFORMATION (continued)

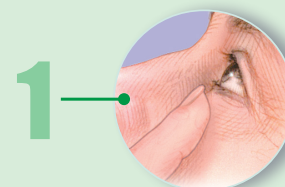
WARNINGS AND PRECAUTIONS

Potential of Vascular Insufficiency: ALPHAGAN® P (brimonidine tartrate ophthalmic solution) 0.1% or 0.15% may potentiate syndromes associated with vascular insufficiency. ALPHAGAN® P should be used with caution in patients with depression, cerebral or coronary insufficiency, Raynaud's phenomenon, orthostatic hypotension, or thromboangiitis obliterans.

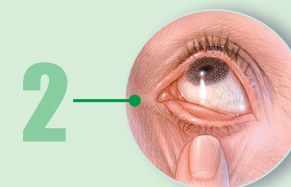
Severe Cardiovascular Disease: Although brimonidine tartrate ophthalmic solution had minimal effect on the blood pressure of patients in clinical studies, caution should be exercised in treating patients with severe cardiovascular disease.

Use ALPHAGAN® P 0.1% the right way

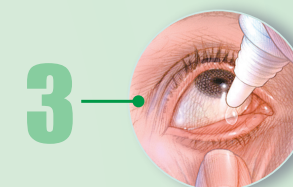
Administering your eyedrops



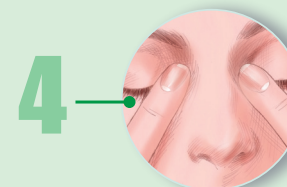
1 — Wash your hands. Tilt your head back so you are looking at the ceiling.*



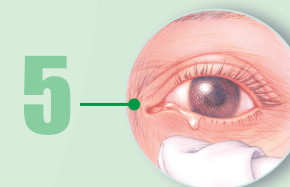
2 — Gently pull down your lower eyelid with your index finger, forming a pocket.



3 — Gently squeeze 1 drop of medication into the pocket. Do not touch the bottle tip to your eye, your fingers, or anything else.



4 — Close your eyes and lightly press your fingers on the inside corners of your eyes.



5 — Carefully blot away any excess liquid that may be on the skin around your eye.

*With certain glaucoma eyedrops, first you will need to remove your contact lenses, if you wear them. Wait 15 minutes after using your eyedrops before you put your contact lenses back into your eyes.

IMPORTANT SAFETY INFORMATION (continued)

WARNINGS AND PRECAUTIONS (continued)

Contamination of Topical Ophthalmic Products After Use: There have been reports of bacterial keratitis associated with the use of multiple-dose containers of topical ophthalmic products. These containers had been inadvertently contaminated by patients who, in most cases, had a concurrent corneal disease or a disruption of the ocular epithelial surface.

Please see additional Important Safety Information on the following pages.

Ensure you get the **GREEN BOTTLE**

Your doctor specifically prescribed **ALPHAGAN® P 0.1%**

- ALPHAGAN® P 0.1% is specifically formulated to contain a lower concentration of brimonidine (the active drug) with the same eye pressure-lowering effects as original ALPHAGAN® 0.2%¹
- In clinical trials, ALPHAGAN® P 0.1% had fewer patients discontinue treatment due to side effects than ALPHAGAN® 0.2% (8.8% of 215 patients compared to 15.1% of 218 patients)¹

IMPORTANT SAFETY INFORMATION (continued)

DRUG INTERACTIONS

Antihypertensives/Cardiac Glycosides: Because ALPHAGAN® P (brimonidine tartrate ophthalmic solution) 0.1% or 0.15% may reduce blood pressure, caution in using drugs such as antihypertensives and/or cardiac glycosides with ALPHAGAN® P is advised.

CNS Depressants: Although specific drug interaction studies have not been conducted with ALPHAGAN® P, the possibility of an additive or potentiating effect with CNS depressants (alcohol, barbiturates, opiates, sedatives, or anesthetics) should be considered.

Check your prescription at the pharmacy to ensure you get the GREEN BOTTLE—ALPHAGAN® P 0.1%

Remember, there is no FDA-approved generic formulation for ALPHAGAN® P 0.1%²

Save with **ALPHAGAN® P 0.1%**

- Visit www.AlphaganP.com for savings up to \$25 and more product information
- Ask your doctor to prescribe the largest covered bottle size. It may save on your monthly co-pays
- Ask your doctor or eye care technician about additional savings programs available at the office



IMPORTANT SAFETY INFORMATION (continued)

DRUG INTERACTIONS (continued)

Tricyclic Antidepressants: Tricyclic antidepressants have been reported to blunt the hypotensive effect of systemic clonidine. It is not known whether the concurrent use of these agents with ALPHAGAN® P (brimonidine tartrate ophthalmic solution) 0.1% or 0.15% in humans can lead to resulting interference with the IOP-lowering effect. Caution is advised in patients taking tricyclic antidepressants, which can affect the metabolism and uptake of circulating amines.

Monoamine Oxidase Inhibitors: Monoamine oxidase (MAO) inhibitors may theoretically interfere with the metabolism of brimonidine and potentially result in an increased systemic side effect such as hypotension. Caution is advised in patients taking MAO inhibitors, which can affect the metabolism and uptake of circulating amines.

Please see additional Important Safety Information on back cover.

Ask your doctor to prescribe the largest bottle size of ALPHAGAN® P 0.1% covered under your insurance plan—you could save on your monthly co-pays

month _____ year _____

	Sunday		Monday		Tuesday	
	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Medication 1 _____ AM ☐ midday ☐ PM ☐

Medication 2 _____ AM ☐ midday ☐ PM ☐

Medication 3 _____ AM ☐ midday ☐ PM ☐

Wednesday		Thursday		Friday		Saturday	
R eye	L eye	R eye	L eye	R eye	L eye	R eye	L eye
Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

month _____ year _____

	Sunday		Monday		Tuesday	
	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Medication 1 _____ AM ☐ midday ☐ PM ☐

Medication 2 _____ AM ☐ midday ☐ PM ☐

Medication 3 _____ AM ☐ midday ☐ PM ☐

	Wednesday		Thursday		Friday		Saturday	
	R eye	L eye	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Please see the Indication and Important Safety Information on pages 1 to 5 and on the back cover.

month _____ year _____

	Sunday		Monday		Tuesday	
	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Medication 1 _____ AM ☐ midday ☐ PM ☐

Medication 2 _____ AM ☐ midday ☐ PM ☐

Medication 3 _____ AM ☐ midday ☐ PM ☐

Wednesday		Thursday		Friday		Saturday	
R eye	L eye	R eye	L eye	R eye	L eye	R eye	L eye
Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

month _____ year _____

	Sunday		Monday		Tuesday	
	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Medication 1 _____ AM ☐ midday ☐ PM ☐

Medication 2 _____ AM ☐ midday ☐ PM ☐

Medication 3 _____ AM ☐ midday ☐ PM ☐

	Wednesday		Thursday		Friday		Saturday	
	R eye	L eye	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Please see the Indication and Important Safety Information on pages 1 to 5 and on the back cover.

month _____ year _____

	Sunday		Monday		Tuesday	
	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Medication 1 _____ AM ☐ midday ☐ PM ☐

Medication 2 _____ AM ☐ midday ☐ PM ☐

Medication 3 _____ AM ☐ midday ☐ PM ☐

	Wednesday		Thursday		Friday		Saturday	
	R eye	L eye	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

month _____ year _____

	Sunday		Monday		Tuesday	
	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Medication 1 _____ AM ☐ midday ☐ PM ☐

Medication 2 _____ AM ☐ midday ☐ PM ☐

Medication 3 _____ AM ☐ midday ☐ PM ☐

	Wednesday		Thursday		Friday		Saturday	
	R eye	L eye	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Please see the Indication and Important Safety Information on pages 1 to 5 and on the back cover.

month _____ year _____

	Sunday		Monday		Tuesday	
	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Medication 1 _____ AM ☐ midday ☐ PM ☐

Medication 2 _____ AM ☐ midday ☐ PM ☐

Medication 3 _____ AM ☐ midday ☐ PM ☐

Wednesday		Thursday		Friday		Saturday	
R eye	L eye	R eye	L eye	R eye	L eye	R eye	L eye
Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

month _____ year _____

	Sunday		Monday		Tuesday	
	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Medication 1 _____ AM ☐ midday ☐ PM ☐

Medication 2 _____ AM ☐ midday ☐ PM ☐

Medication 3 _____ AM ☐ midday ☐ PM ☐

	Wednesday		Thursday		Friday		Saturday	
	R eye	L eye	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Please see the Indication and Important Safety Information on pages 1 to 5 and on the back cover.

month _____ year _____

	Sunday		Monday		Tuesday	
	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Medication 1 _____ AM ☐ midday ☐ PM ☐

Medication 2 _____ AM ☐ midday ☐ PM ☐

Medication 3 _____ AM ☐ midday ☐ PM ☐

Wednesday		Thursday		Friday		Saturday	
R eye	L eye	R eye	L eye	R eye	L eye	R eye	L eye
Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

month _____ year _____

	Sunday		Monday		Tuesday	
	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Medication 1 _____ AM ☐ midday ☐ PM ☐

Medication 2 _____ AM ☐ midday ☐ PM ☐

Medication 3 _____ AM ☐ midday ☐ PM ☐

Wednesday		Thursday		Friday		Saturday	
R eye	L eye	R eye	L eye	R eye	L eye	R eye	L eye
Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Date _____		Date _____		Date _____		Date _____	
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Please see the Indication and Important Safety Information on pages 1 to 5 and on the back cover.

month _____ year _____

	Sunday		Monday		Tuesday	
	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Medication 1 _____ AM ☐ midday ☐ PM ☐
 Medication 2 _____ AM ☐ midday ☐ PM ☐
 Medication 3 _____ AM ☐ midday ☐ PM ☐

	Wednesday		Thursday		Friday		Saturday	
	R eye	L eye	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

month _____ year _____

	Sunday		Monday		Tuesday	
	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____	
Med 1	○○○	○○○	○○○	○○○	○○○	○○○
Med 2	○○○	○○○	○○○	○○○	○○○	○○○
Med 3	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Medication 1 _____ AM ☐ midday ☐ PM ☐

Medication 2 _____ AM ☐ midday ☐ PM ☐

Medication 3 _____ AM ☐ midday ☐ PM ☐

	Wednesday		Thursday		Friday		Saturday	
	R eye	L eye	R eye	L eye	R eye	L eye	R eye	L eye
	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

	Date _____		Date _____		Date _____		Date _____	
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○
	○○○	○○○	○○○	○○○	○○○	○○○	○○○	○○○

Notes _____

Please see the Indication and Important Safety Information on pages 1 to 5 and on the back cover.

This image shows a single sheet of white paper with horizontal blue or grey ruling lines. The lines are evenly spaced and run across the width of the page. There is no handwriting or other markings on the paper.

[illegible]

IMPORTANT SAFETY INFORMATION (continued)

ADVERSE REACTIONS

Adverse reactions occurring in approximately 10% to 20% of the subjects receiving brimonidine ophthalmic solution (0.1% to 0.2%) included: allergic conjunctivitis, conjunctival hyperemia, and eye pruritus. Adverse reactions occurring in approximately 5% to 9% included: burning sensation, conjunctival folliculosis, hypertension, ocular allergic reaction, oral dryness, and visual disturbance.

Please see the Indication and additional Important Safety Information on pages 1 to 5.

Please see accompanying full Prescribing Information.



1. Data on file, Allergan, Inc., 2003; Study number:190342-021. 2. US Food and Drug Administration. Drugs@FDA. US Food and Drug Administration website. <http://www.accessdata.fda.gov/scripts/cder/drugsatfda/>. Accessed September 27, 2013.

Alphagan[®] P 0.1%
(brimonidine tartrate ophthalmic solution) 0.1%

HIGHLIGHTS OF PRESCRIBING INFORMATION

These highlights do not include all the information needed to use **ALPHAGAN® P** safely and effectively. See full prescribing information for **ALPHAGAN® P**.

ALPHAGAN® P (brimonidine tartrate ophthalmic solution) 0.1% and 0.15%
Initial U.S. Approval: 1996

INDICATIONS AND USAGE

ALPHAGAN® P is an alpha adrenergic receptor agonist indicated for the reduction of elevated intraocular pressure (IOP) in patients with open-angle glaucoma or ocular hypertension. (1)

DOSAGE AND ADMINISTRATION

- One drop in the affected eye(s), three times daily, approximately 8 hours apart. (2)

DOSAGE FORMS AND STRENGTHS

- Solution containing 1 or 1.5 mg/mL brimonidine tartrate. (3)

CONTRAINDICATIONS

- Neonates and infants (under the age of 2 years). (4.1)

WARNINGS AND PRECAUTIONS

Potentiation of vascular insufficiency. (5.1)

ADVERSE REACTIONS

Most common adverse reactions occurring in approximately 5% to 20% of patients receiving brimonidine ophthalmic solution (0.1%-0.2%) included allergic conjunctivitis, burning sensation, conjunctival folliculosis, conjunctival hyperemia, eye pruritus, hypertension, ocular allergic reaction, oral dryness, and visual disturbance. (6.1)

To report SUSPECTED ADVERSE REACTIONS, contact Allergan at 1-800-433-8871 or the FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

DRUG INTERACTIONS

- Antihypertensives/cardiac glycosides may lower blood pressure. (7.1)
- Use with CNS depressants may result in an additive or potentiating effect. (7.2)
- Tricyclic antidepressants may potentially blunt the hypotensive effect of systemic clonidine. (7.3)
- Monoamine oxidase inhibitors may result in increased hypotension. (7.4)

USE IN SPECIFIC POPULATIONS

- Use with caution in children ≥ 2 years of age. (8.4)

See 17 for PATIENT COUNSELING INFORMATION.

Revised: 09/2013

FULL PRESCRIBING INFORMATION: CONTENTS*

- 1 INDICATIONS AND USAGE
- 2 DOSAGE AND ADMINISTRATION
- 3 DOSAGE FORMS AND STRENGTHS
- 4 CONTRAINDICATIONS
 - 4.1 Neonates and Infants (under the age of 2 years)
 - 4.2 Hypersensitivity Reactions
- 5 WARNINGS AND PRECAUTIONS
 - 5.1 Potentiation of Vascular Insufficiency
 - 5.2 Severe Cardiovascular Disease
 - 5.3 Contamination of Topical Ophthalmic Products After Use
- 6 ADVERSE REACTIONS
 - 6.1 Clinical Studies Experience
 - 6.2 Postmarketing Experience
- 7 DRUG INTERACTIONS
 - 7.1 Antihypertensives/Cardiac Glycosides
 - 7.2 CNS Depressants
 - 7.3 Tricyclic Antidepressants
 - 7.4 Monoamine Oxidase Inhibitors

- 8 USE IN SPECIFIC POPULATIONS
 - 8.1 Pregnancy
 - 8.3 Nursing Mothers
 - 8.4 Pediatric Use
 - 8.5 Geriatric Use
 - 8.6 Special Populations
- 10 OVERDOSAGE
- 11 DESCRIPTION
- 12 CLINICAL PHARMACOLOGY
 - 12.1 Mechanism of Action
 - 12.3 Pharmacokinetics
- 13 NONCLINICAL TOXICOLOGY
 - 13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility
- 14 CLINICAL STUDIES
- 16 HOW SUPPLIED/STORAGE AND HANDLING
- 17 PATIENT COUNSELING INFORMATION

*Sections or subsections omitted from the full prescribing information are not listed.

FULL PRESCRIBING INFORMATION

1 INDICATIONS AND USAGE

ALPHAGAN® P (brimonidine tartrate ophthalmic solution) 0.1% or 0.15% is an alpha adrenergic receptor agonist indicated for the reduction of elevated intraocular pressure (IOP) in patients with open-angle glaucoma or ocular hypertension.

2 DOSAGE AND ADMINISTRATION

The recommended dose is one drop of **ALPHAGAN® P** in the affected eye(s) three times daily, approximately 8 hours apart. **ALPHAGAN® P** ophthalmic solution may be used concomitantly with other topical ophthalmic drug products to lower intraocular pressure. If more than one topical ophthalmic product is to be used, the different products should be instilled at least 5 minutes apart.

3 DOSAGE FORMS AND STRENGTHS

Solution containing 1 mg/mL or 1.5 mg/mL brimonidine tartrate.

4 CONTRAINDICATIONS

4.1 Neonates and Infants (under the age of 2 years)

ALPHAGAN® P is contraindicated in neonates and infants (under the age of 2 years).

4.2 Hypersensitivity Reactions

ALPHAGAN® P is contraindicated in patients who have exhibited a hypersensitivity reaction to any component of this medication in the past.

5 WARNINGS AND PRECAUTIONS

5.1 Potentiation of Vascular Insufficiency

ALPHAGAN® P may potentiate syndromes associated with vascular insufficiency. **ALPHAGAN® P** should be used with caution in patients with depression, cerebral or coronary insufficiency, Raynaud's phenomenon, orthostatic hypotension, or thromboangiitis obliterans.

5.2 Severe Cardiovascular Disease

Although brimonidine tartrate ophthalmic solution had minimal effect on the blood pressure of patients in clinical studies, caution should be exercised in treating patients with severe cardiovascular disease.

5.3 Contamination of Topical Ophthalmic Products After Use

There have been reports of bacterial keratitis associated with the use of multiple-dose containers of topical ophthalmic products. These containers had been inadvertently contaminated by patients who, in most cases, had a concurrent corneal disease or a disruption of the ocular epithelial surface (see **PATIENT COUNSELING INFORMATION**, 17).

6 ADVERSE REACTIONS

6.1 Clinical Studies Experience

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.

Adverse reactions occurring in approximately 10-20% of the subjects receiving brimonidine ophthalmic solution (0.1-0.2%) included: allergic conjunctivitis, conjunctival hyperemia, and eye pruritus. Adverse reactions occurring in approximately 5-9% included: burning sensation, conjunctival folliculosis, hypertension, ocular allergic reaction, oral dryness, and visual disturbance.

Adverse reactions occurring in approximately 1-4% of the subjects receiving brimonidine ophthalmic solution (0.1-0.2%) included: abnormal taste, allergic reaction, asthenia, blepharitis, blepharoconjunctivitis, blurred vision, bronchitis, cataract, conjunctival edema, conjunctival hemorrhage, conjunctivitis, cough, dizziness, dyspepsia, dyspnea, epiphora, eye discharge, eye dryness, eye irritation, eye pain, eyelid edema, eyelid erythema, fatigue, flu syndrome, follicular conjunctivitis, foreign body sensation, gastrointestinal disorder, headache, hypercholesterolemia, hypotension, infection (primarily colds and respiratory infections), insomnia, keratitis, lid disorder, pharyngitis, photophobia, rash, rhinitis, sinus infection, sinusitis, somnolence, stinging, superficial punctate keratopathy, tearing, visual field defect, vitreous detachment, vitreous disorder, vitreous floaters, and worsened visual acuity.

The following reactions were reported in less than 1% of subjects: corneal erosion, hordeolum, nasal dryness, and taste perversion.

6.2 Postmarketing Experience

The following reactions have been identified during postmarketing use of brimonidine tartrate ophthalmic solutions in clinical practice. Because they are reported voluntarily from a population of unknown size, estimates of frequency cannot be made. The reactions, which have been chosen for inclusion due to either their seriousness, frequency of reporting, possible causal connection to brimonidine tartrate ophthalmic solutions, or a combination of these factors, include: bradycardia, depression, hypersensitivity, iritis, keratoconjunctivitis sicca, miosis, nausea, skin reactions (including erythema, eyelid pruritus, rash, and vasodilation), syncope, and tachycardia. Apnea, bradycardia, coma, hypotension, hypothermia, hypotonia, lethargy, pallor, respiratory depression, and somnolence have been reported in infants receiving brimonidine tartrate ophthalmic solutions.

7 DRUG INTERACTIONS

7.1 Antihypertensives/Cardiac Glycosides

Because **ALPHAGAN® P** may reduce blood pressure, caution in using drugs such as antihypertensives and/or cardiac glycosides with **ALPHAGAN® P** is advised.

7.2 CNS Depressants

Although specific drug interaction studies have not been conducted with **ALPHAGAN® P**, the possibility of an additive or potentiating effect with CNS depressants (alcohol, barbiturates, opiates, sedatives, or anesthetics) should be considered.

7.3 Tricyclic Antidepressants

Tricyclic antidepressants have been reported to blunt the hypotensive effect of systemic clonidine. It is not known whether the concurrent use of these agents with **ALPHAGAN® P** in humans can lead to resulting interference with the IOP lowering effect. Caution is advised in patients taking tricyclic antidepressants which can affect the metabolism and uptake of circulating amines.

7.4 Monoamine Oxidase Inhibitors

Monoamine oxidase (MAO) inhibitors may theoretically interfere with the metabolism of brimonidine and potentially result in an increased systemic side-effect such as hypotension. Caution is advised in patients taking MAO inhibitors which can affect the metabolism and uptake of circulating amines.

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Pregnancy Category B: Teratogenicity studies have been performed in animals.

Brimonidine tartrate was not teratogenic when given orally during gestation days 6 through 15 in rats and days 6 through 18 in rabbits. The highest doses of brimonidine tartrate in rats (2.5 mg/kg/day) and rabbits (5.0 mg/kg/day) achieved AUC exposure values 360- and 20-fold higher, or 260- and 15-fold higher, respectively, than similar values estimated in humans treated with **ALPHAGAN® P** 0.1% or 0.15%, 1 drop in both eyes three times daily.

There are no adequate and well-controlled studies in pregnant women; however, in animal studies, brimonidine crossed the placenta and entered into the fetal circulation to a limited extent. Because animal reproduction studies are not always predictive of human response, **ALPHAGAN® P** should be used during pregnancy only if the potential benefit to the mother justifies the potential risk to the fetus.

8.3 Nursing Mothers

It is not known whether brimonidine tartrate is excreted in human milk, although in animal studies, brimonidine tartrate has been shown to be excreted in breast milk. Because of the potential for serious adverse reactions from **ALPHAGAN® P** in nursing infants, a decision should be made whether to discontinue nursing or to discontinue the drug, taking into account the importance of the drug to the mother.

8.4 Pediatric Use

ALPHAGAN® P is contraindicated in children under the age of 2 years (see **CONTRAINDICATIONS**, 4.1). During postmarketing surveillance, apnea, bradycardia, coma, hypotension, hypothermia, hypotonia, lethargy, pallor, respiratory depression, and somnolence have been reported in infants receiving brimonidine. The safety and effectiveness of brimonidine tartrate have not been studied in children below the age of 2 years.

In a well-controlled clinical study conducted in pediatric glaucoma patients (ages 2 to 7 years) the most commonly observed adverse reactions with brimonidine tartrate ophthalmic solution 0.2% dosed three times daily were somnolence (50-83% in patients ages 2 to 6 years) and decreased alertness. In pediatric patients 7 years of age (>20 kg), somnolence appears to occur less frequently (25%). Approximately 16% of patients on brimonidine tartrate ophthalmic solution discontinued from the study due to somnolence.

8.5 Geriatric Use

No overall differences in safety or effectiveness have been observed between elderly and other adult patients.

8.6 Special Populations

ALPHAGAN® P has not been studied in patients with hepatic impairment.

ALPHAGAN® P has not been studied in patients with renal impairment. The effect of dialysis on brimonidine pharmacokinetics in patients with renal failure is not known.

10 OVERDOSAGE

Very limited information exists on accidental ingestion of brimonidine in adults; the only adverse reaction reported to date has been hypotension. Symptoms of brimonidine overdose have been reported in neonates, infants, and children receiving **ALPHAGAN® P** as part of medical treatment of congenital glaucoma or by accidental oral ingestion (see **USE IN SPECIFIC POPULATIONS**, 8.4). Treatment of an oral overdose includes supportive and symptomatic therapy; a patent airway should be maintained.

11 DESCRIPTION

ALPHAGAN® P (brimonidine tartrate ophthalmic solution) 0.1% or 0.15%, sterile, is a relatively selective alpha-2 adrenergic receptor agonist (topical intraocular pressure lowering agent).

The structural formula of brimonidine tartrate is:

5-Bromo-6-(2-imidazolidinylideneamino) quinoxaline L-tartrate; MW= 442.24

In solution, **ALPHAGAN® P** (brimonidine tartrate ophthalmic solution) has a clear, greenish-yellow color. It has an osmolality of 250-350 mOsmol/kg and a pH of 7.4-8.0 (0.1%) or 6.9-7.4 (0.15%).

Brimonidine tartrate appears as an off-white to pale-yellow powder and is soluble in both water (0.6 mg/mL) and in the product vehicle (1.4 mg/mL) at pH 7.7.

Each mL of **ALPHAGAN® P** contains the active ingredient brimonidine tartrate 0.1% (1 mg/mL) or 0.15% (1.5 mg/mL) with the inactive ingredients sodium carboxymethylcellulose; sodium borate; boric acid; sodium chloride; potassium chloride; calcium chloride; magnesium chloride; PURITE® 0.005% (0.05 mg/mL) as a preservative; purified water; and hydrochloric acid and/or sodium hydroxide to adjust pH.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

ALPHAGAN® P is a relatively selective alpha-2 adrenergic receptor agonist with a peak ocular hypotensive effect occurring at two hours post-dosing.

Fluorophotometric studies in animals and humans suggest that brimonidine tartrate has a dual mechanism of action by reducing aqueous humor production and increasing uveoscleral outflow.

12.3 Pharmacokinetics

Absorption

After ocular administration of either a 0.1% or 0.2% solution, plasma concentrations peaked within 0.5 to 2.5 hours and declined with a systemic half-life of approximately 2 hours.

Distribution

The protein binding of brimonidine has not been studied.

Metabolism

In humans, brimonidine is extensively metabolized by the liver.

Excretion

Urinary excretion is the major route of elimination of brimonidine and its metabolites. Approximately 87% of an orally-administered radioactive dose of brimonidine was eliminated within 120 hours, with 74% found in the urine.

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

No compound-related carcinogenic effects were observed in either mice or rats following a 21-month and 24-month study, respectively. In these studies, dietary administration of brimonidine tartrate at doses up to 2.5 mg/kg/day in mice and 1 mg/kg/day in rats achieved 150 and 120 times or 90 and 80 times, respectively, the plasma C_{max} drug concentration in humans treated with one drop of **ALPHAGAN® P** 0.1% or 0.15% into both eyes 3 times per day, the recommended daily human dose.

Brimonidine tartrate was not mutagenic or clastogenic in a series of *in vitro* and *in vivo* studies including the Ames bacterial reversion test, chromosomal aberration assay in Chinese Hamster Ovary (CHO) cells, and three *in vivo* studies in CD-1 mice: a host-mediated assay, cytogenetic study, and dominant lethal assay.

Reproduction and fertility studies in rats with brimonidine tartrate demonstrated no adverse effect on male or female fertility at doses which achieve up to approximately 125 and 90 times the systemic exposure following the maximum recommended human ophthalmic dose of **ALPHAGAN® P** 0.1% or 0.15%, respectively.

14 CLINICAL STUDIES

Elevated IOP presents a major risk factor in glaucomatous field loss. The higher the level of IOP, the greater the likelihood of optic nerve damage and visual field loss. Brimonidine tartrate has the action of lowering intraocular pressure with minimal effect on cardiovascular and pulmonary parameters.

Clinical studies were conducted to evaluate the safety, efficacy, and acceptability of **ALPHAGAN® P** (brimonidine tartrate ophthalmic solution) 0.15% compared with **ALPHAGAN®** administered three-times-daily in patients with open-angle glaucoma or ocular hypertension. Those results indicated that **ALPHAGAN® P** (brimonidine tartrate ophthalmic solution) 0.15% is comparable in IOP lowering effect to **ALPHAGAN®** (brimonidine tartrate ophthalmic solution) 0.2%, and effectively lowers IOP in patients with open-angle glaucoma or ocular hypertension by approximately 2-6 mmHg.

A clinical study was conducted to evaluate the safety, efficacy, and acceptability of **ALPHAGAN® P** (brimonidine tartrate ophthalmic solution) 0.1% compared with **ALPHAGAN®** administered three-times-daily in patients with open-angle glaucoma or ocular hypertension. Those results indicated that **ALPHAGAN® P** (brimonidine tartrate ophthalmic solution) 0.1% is equivalent in IOP lowering effect to **ALPHAGAN®** (brimonidine tartrate ophthalmic solution) 0.2%, and effectively lowers IOP in patients with open-angle glaucoma or ocular hypertension by approximately 2-6 mmHg.

16 HOW SUPPLIED/STORAGE AND HANDLING

ALPHAGAN® P is supplied sterile, in teal opaque plastic LDPE bottles and tips, with purple high impact polystyrene (HIPS) caps as follows:

0.1%	5 mL in 10 mL bottle	NDC 0023-9321-05
	10 mL in 10 mL bottle	NDC 0023-9321-10
	15 mL in 15 mL bottle	NDC 0023-9321-15

0.15%	5 mL in 10 mL bottle	NDC 0023-9177-05
	10 mL in 10 mL bottle	NDC 0023-9177-10
	15 mL in 15 mL bottle	NDC 0023-9177-15

Storage: Store at 15°-25°C (59°-77°F).

17 PATIENT COUNSELING INFORMATION

Patients should be instructed that ocular solutions, if handled improperly or if the tip of the dispensing container contacts the eye or surrounding structures, can become contaminated by common bacteria known to cause ocular infections. Serious damage to the eye and subsequent loss of vision may result from using contaminated solutions (see **WARNINGS AND PRECAUTIONS**, 5.3). Always replace the cap after using. If solution changes color or becomes cloudy, do not use. Do not use the product after the expiration date marked on the bottle.

Patients also should be advised that if they have ocular surgery or develop an intercurrent ocular condition (e.g., trauma or infection), they should immediately seek their physician's advice concerning the continued use of the present multidose container.

If more than one topical ophthalmic drug is being used, the drugs should be administered at least five minutes apart.

As with other similar medications, **ALPHAGAN® P** may cause fatigue and/or drowsiness in some patients. Patients who engage in hazardous activities should be cautioned of the potential for a decrease in mental alertness.

©2014 Allergan, Inc.
Irvine, CA 92612, U.S.A.
® marks owned by Allergan, Inc.

Patented. See: www.allergan.com/products/patent_notices

Made in the U.S.A.

Based on 71816US16

APC630I14