

## Uptravi - (0.2, 0.4, 0.6, 0.8, 1, 1.2, 1.4, 1.6 mg; Tablet,)

<b>Generic Name</b>	Selexipag	<b>Innovator</b>	Actelion pharmaceuticals
<b>Dosage</b>	0.2, 0.4, 0.6, 0.8, 1, 1.2, 1.4, 1.6 mg; Tablet,	<b>Branded US Sales</b>	More Than \$1000 mn
<b>Probable FTF</b>	Less Than 5	<b>Known Para IV Filers</b>	More Than 5
<b>Other ANDA developers</b>	More Than 5	<b>Tentative Approvals</b>	Less Than 5
<b>Final Approvals</b>	None	<b>Generic Launches</b>	None
<b>Indication</b>	UPTRAVI is indicated for the treatment of pulmonary arterial hypertension (PAH, WHO Group I) to delay disease progression and reduce the risk of hospitalization for PAH.		
<b>Complexities</b>	Yes		

### Chronology Of Events

Please Contact [contact@researchdelta.com](mailto:contact@researchdelta.com) to get Detailed Information.

### Executive Summary

Please Contact [contact@researchdelta.com](mailto:contact@researchdelta.com) to get Detailed Information.

### Patent Status

Please Contact [contact@researchdelta.com](mailto:contact@researchdelta.com) to get Detailed Information.

### Launch Timelines and Competition

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## Uptravi - (1.8 mg/vial ; Powder, Intravenous)

<b>Generic Name</b>	Selexipag	<b>Innovator</b>	ACTELION
<b>Dosage</b>	1.8 mg/vial ; Powder, Intravenous	<b>Branded US Sales</b>	More Than \$1000 mn
<b>Probable FTF</b>	Less Than 5	<b>Known Para IV Filers</b>	Less Than 5
<b>Other ANDA developers</b>	None	<b>Tentative Approvals</b>	Less Than 5
<b>Final Approvals</b>	None	<b>Generic Launches</b>	None
<b>Indication</b>	UPTRAVI is a prostacyclin receptor agonist indicated for the treatment of pulmonary arterial hypertension (PAH, WHO Group I) to delay disease progression and reduce the risk of hospitalization for PAH.		
<b>Complexities</b>	Yes		

### Chronology Of Events

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