#### VI.2 Elements for a Public Summary

#### VI.2.1 Overview of disease epidemiology

During anaesthesia the patient undergoing anesthesia may experience a reduction in normal blood pressure (typically  $\geq$  20% decrease from normal blood pressure), i.e. hypotension. It is estimated that one-third of patients undergoing spinal anesthesia experience hypotension. This hypotension during spinal anaesthesia is considered to be due to decrease in amount of blood returned to the heart and due to reduction in amount of blood pumped out of heart. Hypotension may also occur after general anesthesia. Prolonged hypotension during anesthesia is associated with increased mortality and also unwanted side effects related to both heart and other bodily systems. Therefore, during the surgery restoration of blood pressure is the immediate requirement. Increased or normalised BP is the appropriate endpoint for a drug such as phenylephrine which acts by tightening blood vessels. Also, hypotension is one of the most important causes of nausea and vomiting particularly in the

initial period after initiation of spinal anesthesia.

### VI.2.2 Summary of treatment benefits

Phenylephrine is a very selective chemical agent which specifically binds to certain protein to increase the blood pressure when it is lower than normal blood pressure. It has the advantage of not being an agent which alters force of muscle contractions or heart rate. It strictly elevates the blood pressure without increasing the heart rate or contractility. This is especially useful if the heart is already prone to other heart muscle diseases or increase in heart rate. Phenylephrine injection has been used in different medical settings, notably in critical care, cardiology, and anesthesia for over 75 years. Currently, phenylephrine is one of the most common drug used to elevate the blood pressure during the care of women during and after pregnancy.

### VI.2.3 Unknowns relating to treatment benefits

There are no adequate and well-controlled studies in use of phenylephrine in pregnant women. Limited data do not indicate any increased risk of birth defect malformations. Animal studies are insufficient with respect to effects on pregnancy, fetal development, parturition or postnatal development. The potential risk for humans is unknown. Therefore, phenylephrine

should not be used during pregnancy unless clearly necessary.

# VI.2.4 Summary of safety concerns

# Important identified risks

Risk	What is known	Preventability
Arterial hypertension	In the literature arterial hypertension is reportedly associated with phenylephrine.	Hypertension due to phenylephrine can be prevented by careful selection of dose and by monitoring of blood pressure during treatment.
Arrhythmia	Phenylephrine is known to cause arrhythmia	Warning about this risk and increased risk in patients with bradycardia and cardiac diseases are included in the SmPC and PIL.

Risk	What is known (Including reason why it is
	considered a potential risk)
Extravasation	Patients treated with medications that constrict muscles in the blood vessels may be at an increased risk of severe tissue damage if they extravasate. For safe use Fenylefrin Abcur should only be given by a healthcare professional with appropriate training and experience.

Risk	What is known
Use during pregnancy	There is no adequate data from clinical trials for use
	of
	phenylephrine in pregnant women. Animal studies
	are
	insufficient with respect to effects on pregnancy,
	fetal
	development, parturition or postnatal development.
	The
	potential risk for humans is unknown.
	Phenylephrine should not be used during pregnancy
	unless
	clearly necessary.

# VI.2.5 Summary of risk minimisation measures by safety concern

All medicines have a Summary of Product Characteristics (SmPC) which provides physicians,

pharmacists and other health care professionals with details on how to use the medicine, the risks and recommendations for minimising them. An abbreviated version of this in lay language is provided in the form of the package leaflet (PL). The measures in these documents are known as routine risk minimisation measures.

The Summary of Product Characteristics and the Package leaflet for phenylephrine can be found in the phenylephrine's EPAR page.

This medicine has no additional risk minimisation measures.

### VI.2.6 Planned post authorisation development plan

NA

### VI.2.7 Summary of changes to the Risk Management Plan over time

NA