

# OCULAR SURFACE PERMANENCE AND TOXICITY STUDIES OF TACROLIMUS-(HYDROXYPROPYL- $\beta$ -CYCLODEXTRIN) EYEDROPS

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## INTRODUCTION

Noninfectious uveitis is a disease often caused by an autoimmune response, inflammatory cytokines promote the activation of T-cells and trigger recruitment of large numbers of circulation inflammatory leukocytes into the eye. This process may cause irreversible tissue damage and visual impairment. Since tacrolimus inhibits T-cell proliferation and suppresses the release of inflammatory cytokines, it can theoretically be used to reduce inflammatory activity in uveitis patients [1]. Hospital pharmacy prepares tacrolimus eye drops reformulating from parenteral drugs (Prograf<sup>®</sup>) as magistral formulations because there is not any commercial alternative. However, Prograf<sup>®</sup> (tacrolimus solubilized in ethanol) has some irritating compounds that cause discomfort and unpleasantness to the patient, so these excipients had to be removed. Due to tacrolimus poorly solubility, the purpose of this work was to improve the drug solubility complexing tacrolimus with HP $\beta$ CD, evaluate the safety, and study the ocular permanence of the eye drops.

SOLUBILITY	OCULAR PERMANENCE	HETCAM	BCOP

## RESULTS

SOLUBILITY	OCULAR PERMANENCE	HETCAM	BCOP

## CONCLUSIONS

Results reveal tacrolimus solubility improvement and irritation absence. Higher permanence on the ocular surface is achieved with higher HP $\beta$ CD concentrations. Liquifilm<sup>®</sup> eyedrops present less ocular clearance than BSS<sup>®</sup> ones. These formulations would enhance the patient's adherence-to-treatment, reducing eye discomfort.

## REFERENCES

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