Report to the European Commission<sup>1</sup> on Pharmacovigilance audits Carried out in the Agency for Medicinal Products and Medical Devices (HALMED), Croatia

from 12/09/2019 to 10/09/2021

<sup>&</sup>lt;sup>1</sup> Delete as necessary – National Competent Authorities are required to report to the European Commission (Directive 2001/83/EC Art.101(2), European Medicines Agency is required to report to its Management Board (Regulation (EC) No.726/2004 Art 28f)

### 1. INTRODUCTION

The report is prepared in the context of the obligation under Article 101 (2) of the Directive 2001/83/EC states: "Member States shall perform a regular audit of their Pharmacovigilance system and report the results to the Commission on 21 September 2013 at the latest and then every 2 years thereafter."

This report provides an overview of the audit activities conducted from 12/09/2019 to 10/09/2021 by the Agency for Medicinal Products and Medical Devices (hereinafter referred as "HALMED"), Croatia, based on the internal audit which planned to fulfil the Union legal requirements in the field of pharmacovigilance.

For this purpose, HALMED planned and performed 4 internal audits, to check the compatibility of all requirements set in Good pharmacovigilance practices guidelines within HALMED's units responsible for pharmacovigilance activities.

The realisation of achieved goals has been examined during the internal audit whose details are given in the report below.

# 2. DEVELOPMENTS IN THE PHARMACOVIGILANCE SYSTEM SINCE THE LAST REPORT

Significant changes during the reporting period

- Legislation and regulatory
  - No change
- Standards and Procedures
  - No change
- Quality system for Pharmacovigilance Activities
  - Revision of corresponding SOPs connected to GVP Modules that had been revised;
  - Revision of Quality Manual of HALMED (integration of all Quality manuals of specific departments and processes into one Quality Manual)
- Critical Pharmacovigilance Processes
  - The upgrade to the existing databases.
- Other changes
  - The Inspectorate changed name to Inspections Department"
  - The Head of Inspections Department has been changed.
  - The Head of Department for Pharmacovigilance and Rational Pharmacotherapy has been changed.

# 3. INTERNAL AUDIT ACTIVITY FOR THE PERIOD UNDER REVIEW

### 3.1 RISK ASSESSMENT

A risk assessment exercise was conducted in order to determine the pharmacovigilance system audit priorities for the period under review. The final audit strategy was prepared, based on this risk assessment, and was approved by the quality manager of HALMED.

### 3.2 SUMMARY OF THE AUDITS FOR THE PERIOD UNDER REVIEW

# 3.2.1 AUDIT ASSIGNMENTS FOR THE PERIOD UNDER REVIEW

All listed audits are performed in line with the guidance provided in the GVP Module IV Pharmacovigilance audits.

Audit No	Audit title	Date of audit report
3-20	Internal audit 03/20	23.09.2020.
16-20	Internal audit 16/20	01.12.2020.
2-21	Internal audit 02/21	17.03.2021.
8-21	Internal audit 08/21	09.09.2021

### 3.2.2 AUDIT REPORT

### 3.2.2.1 Objective and scope

The objective of the audits was to determine the compliance of HALMED pharmacovigilance system with the Guideline on good pharmacovigilance practices.

The scope of the audit included compliance with stated Modules of the Guideline on good pharmacovigilance practices.

### Internal audit 03/20

- Module V
- Module VIII
- Module IX
- Module XVI

### Internal audit 16/20

Module III

### Internal audit 02/21

- Module II
- Module X
- Module XV

### Internal audit 08/21

- Module I
- Module IV
- Module VI
- Module VII

# 3.2.2.2 Audit body

The Office for quality management of HALMED

# 3.2.2.3 Opinion

Sufficient and appropriate audit procedure has been conducted and evidence gathered to support the accuracy of the conclusions reached and contained in this report. The conclusions were based on a comparison of the situations as they existed at the time of audit against the audit criteria defined in the Quality manual and corresponding standard operating procedures.

The internal auditing team concluded that no critical nonconformities were noted as defined by GVP Module IV (see 3.2.2.4).

Evaluated pharmacovigilance processes are adequate, appropriate, and effective to provide reasonable assurance that potential risks for public health are being properly managed and that the objectives are met.

Despite some difficulties noted, the overall opinion for the reported period on the HALMED Pharmacovigilance quality system is that the system functioned very well and it can assure the high quality of pharmacovigilance activities.

3.2.2.4 Audit outcomes and actions

In 4 audit no outcomes are reported/rated as 'Critical' or 'Major', in line with the guidance provided in the GVP Module IV Pharmacovigilance audits

Audit No	Find No	Audit No Find No Audit outcomes description	Grading	Action short description	Action end date	Comments on status of actions	Type of follow- up required
1	1	1	1	l.	1	1	
1	1		ı	ı	1	-	ı

# 3.2.2.5 Summary of action plan for current reporting period

The following table provides an overview of audit outcomes and their implementation.

For action from audit outcome	Total	Number implemented	Number not implemented	ented
graded as:			Not started	In progress
Critical	0	0	0	0
Major	0	0	0	0
Total	0	0	0	0

# 4. FOLLOW-UP

## 4.1 SUMMARY OF ACTION PLANS FROM PRIOR BIENNIAL REPORTS

The following table provides an overview of earlier audit outcomes issued by the HALMED and their implementation by the HALMED at the period from 12/09/2019 to 10/09/2021

For action from audit outcome graded as:	Total	Number implemented	Number not implemented	
graded as.			Not started	In progress
Critical	0	0	0	0
Major	2	2	0	0
Total	2	2	0	0

There were two ongoing issues in the prior biennial report that have been solved.

### 4.2 OUTSTANDING ISSUES FROM PRIOR BIENNIAL REPORTS

There are no outstanding issues from the last biennial report.

### 5. DECLARATION

The Agency for Medicinal Products and Medical Devices (HALMED) confirms that this report contains a complete account of all pharmacovigilance system audit activity performed in the period under over the contains of our organisation under Directive 2001/83/EC<sup>2</sup>.

Siniša Tomić, PhD Head of Agency

Date 10/09/2021

<sup>2</sup> Delete as necessary – National Competent Authorities are required to perform a regular audit of their Pharmacovigilance system and report the results to the Commission on 21 September 2013 at the latest and then every 2 years thereafter. (Directive 2001/83/EC Art.101(2).

The European Medicines Agency is required regular independent audits of its pharmacovigilance tasks and report the results to its Management Board on a 2-yearly basis. (Regulation (EC) No.726/2004 Art 28f)