

## Portfolio Roadmap 2025-2027



## 2025 and beyond: Network Portfolio roadmap 2025 2026 **Value Stream Product** H+V MAA H Variations form Vet Variat./Renewals..... Product User Interface PUI Product UI (write capabilities) Initials mark, authorisation H+V, Medicines for all & Ancillary Substances **RPM** SIAMED decommissioning SIAMED decommissioning eSub/EURS eCTD v4.0 / EURS Next Central Repository **EMWP EMWP** European Medicines Web Portal PLM Union Product Database Enhancements UPD Enhancements **UPD** Enhancements ePI ePI MVP ePI implementation PMS API (ISO IDMP standard) PMS API: full write capabilities & FHIR upgrade NCA Product Data Upload Interface FHIR Adaptor **PMS** xEVMPD decommissioning xEVMPD decommissioning xEVMPD Replacement Pre-Auth, HTA Reg. & SME Database Pedra decommissioning **RPM** RPM for: Innovation Task Force, Orphan Des., Scientific Advice H+V Scientific Explorer Scientific Explorer II AI/DA Knowledge mining **Knowledge Mining** RW Metadata & Studies catalog (website+API+data population) R&D Data Analytics Platform Clinical Trials Information System **CTIS** Clinical Trials Information System Modernisation SMS Portal SMS EU-SRS Run ESMP ESMP V2.0 **ESMP** ESMP Interoperability **EudraGMDI** Run EudraGMDP EudraGMDP Re-factoring EV Human Strategy Run EV Human MON Run INSP/PD INSP/PD Run UPhV ASU Run ASU Run CMDS Major production go-live

To be PB approved not prioritized vet

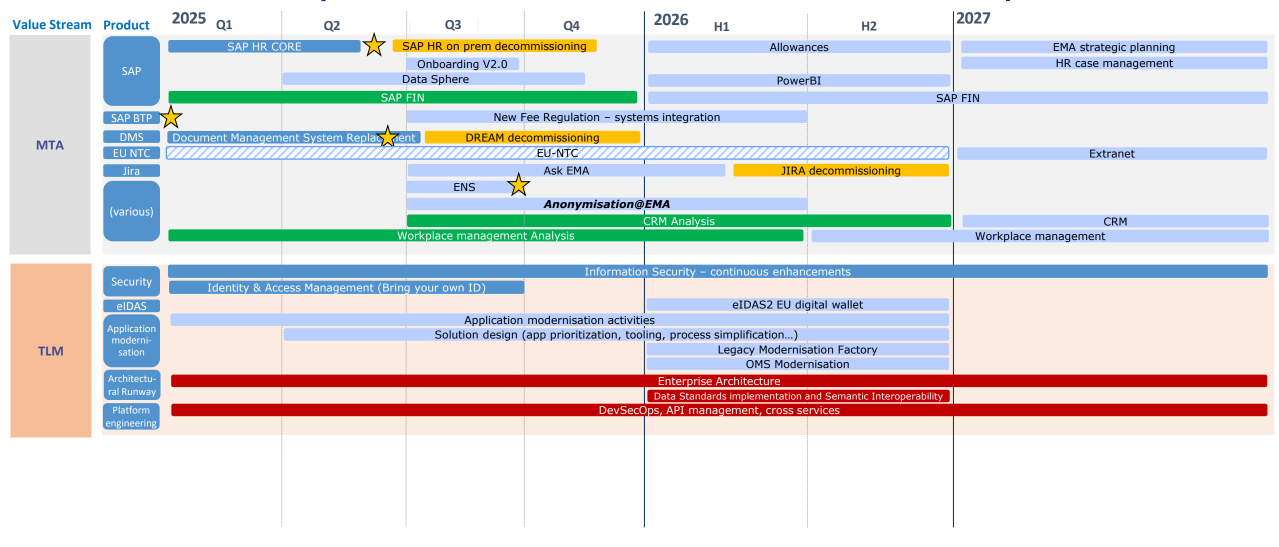
Prioritised

**Analysis** 

**Enabler Epic** 

Run the Business

## 2025 and beyond: Network Portfolio roadmap





## Glossary

Acronym	Name Name
API	Application programming interface
ACT EU	Accelerating Clinical Trials in the EU
Art. 57	Article 57 of Regulation (EU) 726/2004, which requires marketing authorisation holders to electronically submit to the Agency information on all medicinal products for human use authorised in the EU
ASU	Antimicrobials sales and use platform (veterinary products)
AWS	Amazon Web Services
Business Epic	Business epics are large initiatives that deliver Solutions needed by the business/customers
CAP	Centrally authorised product
CDP	Clinical data publication
CMDS	Critical medical devices shortages
CRM	Customer relationship management system
CTIS	Clinical Trials Information System
DA	Data Analytics
DADI	Digital Application Dataset Integration
DARWIN	Data Analysis and Real-World Interrogation Network (DARWIN EU)
DC 2.0	Data Centre 2.0 (cloud-based data centre)
DCP	De-centralised Procedure
DIGIT	DG DIGIT, European Commission Directorate General for Informatics
DREAM	Documents and records management system used at EMA



Acronym	Name
eAF	Electronic application form
ECD	Eudra Common Directory, a directory of individuals and organisations relevant to the operation of the Agency
eCTD	Common Technical Document in electronic format
eIDAS 2.0	European Digital Identity Regulation (Regulation (EU) 2024/1183)
EMRN	European medicines regulatory network
EMWP	European Medicines Web Portal
Enabler Epic	Enabler epics are pieces of work that extend the architectural infrastructure of the solution under development or improve the performance of the value stream
ENS	Early Notification System
ePI	Electronic product information
Epic	An epic is a container with one common objective, for a development initiative large enough to require analysis, definition of a minimal viable product (MVP) and financial approval before implementation. An epic usually takes more than one Programme Increment to complete and is broken into multiple Features.
ESMP	European Medicines Shortages Monitoring Platform
ESMDP	European Medicinal Devices Shortages Monitoring Platform
ESVAC	European Surveillance of Veterinary Antimicrobial Consumption
ETF	Emergency Task Force
EURS	European review system for eCTDs
EU NTC	EU Network Training Centre
EU-SRS	European Substance Reference System
EVVet3	Eudra Vigilance Veterinary version 3 (see UPhV)
EXPAMED	Expert Panels on Medical Devices and In Vitro Diagnostic Devices



HERA Emergency Preparedness and Response Authority HMA Heads of Medicines Agencies IAM Identity and access management IRIS Integrated Regulatory Information System, a secure online platform for handling product-related scientific and regulatory procedures with iSPOC Industry Single Point of Contact ITF Innovation Task Force Jira Service desk ticketing system used at EMA LRSM (Data) Lifecycle Regulatory Submissions Metadata MAA Marketing authorisation application MAH Marketing authorisation holder MD Medical devices MON VS Monitoring Value Stream MRP Mutual Recognition Procedure MSSG Executive Steering Group on Shortages and Safety of Medicinal Products	Acronym	Name
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HMA Heads of Medicines Agencies  IAM Identity and access management  IRIS Integrated Regulatory Information System, a secure online platform for handling product-related scientific and regulatory procedures with iSPOC Industry Single Point of Contact  ITF Innovation Task Force  Jira Service desk ticketing system used at EMA  LRSM (Data) Lifecycle Regulatory Submissions Metadata  MAA Marketing authorisation application  MAH Marketing authorisation holder  MD Medical devices  MON VS Monitoring Value Stream  MRP Mutual Recognition Procedure  MSSG Executive Steering Group on Shortages and Safety of Medicinal Products	IDMP	The ISO IDMP standards specify the use of standardised definitions for the identification and description of medicinal products for human use
IAM Identity and access management IRIS Integrated Regulatory Information System, a secure online platform for handling product-related scientific and regulatory procedures with ISPOC Industry Single Point of Contact ITF Innovation Task Force Jira Service desk ticketing system used at EMA LRSM (Data) Lifecycle Regulatory Submissions Metadata MAA Marketing authorisation application MAH Marketing authorisation holder MD Medical devices MON VS Monitoring Value Stream MRP Mutual Recognition Procedure MSSG Executive Steering Group on Shortages and Safety of Medicinal Products	HERA	Emergency Preparedness and Response Authority
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ISPOC Industry Single Point of Contact ITF Innovation Task Force Jira Service desk ticketing system used at EMA LRSM (Data) Lifecycle Regulatory Submissions Metadata MAA Marketing authorisation application MAH Marketing authorisation holder MD Medical devices MON VS Monitoring Value Stream MRP Mutual Recognition Procedure MSSG Executive Steering Group on Shortages and Safety of Medicinal Products	IAM	Identity and access management
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MSSG Executive Steering Group on Shortages and Safety of Medicinal Products	MON VS	Monitoring Value Stream
	MRP	Mutual Recognition Procedure
MTA VS Managing the Agency Value Stream	MSSG	Executive Steering Group on Shortages and Safety of Medicinal Products
	MTA VS	Managing the Agency Value Stream
MVP Minimum viable product	MVP	Minimum viable product



Acronym	Name Name
NAP	Nationally authorised product
NCA	National Competent Authority
NFR	New fee regulation, Regulation (EU) 2024/568 of 7 February 2024 on fees and charges payable to the European Medicines Agency
NICTAC	Network ICT Advisory Committee represents the network IT community
NPAG	Network Portfolio Advisory Group represents EMA Management Board and Heads of Medicines Agencies (HMA)
NTC	Network Training Centre
P3i	Waterfall project management methodology, used at EMA prior to SAFe Agile
PedRA	Paediatric Records Application, Paediatric database
PAM	Post-authorisation measure
РВ	Portfolio Board
PD	Parallel distribution
PHE / ME	Public health emergency / major events
PI	Planning interval, a three-month period of work
PI ceremony	A quarterly event to plan work for the value stream in the next quarter, ensuring that teams and stakeholders have a shared mission and vision
PLM VS	Product Lifecycle Management Value Stream
PMS	Product (Data) Management Service
PPMT	Public procurement management tool
PRIME	PRIority MEdicines
PO	Product owner (PO) is the agile team member primarily responsible for maximizing the value delivered by the team by ensuring that the team backlog is aligned with customer and stakeholder needs.
PSUSA	PSUR single assessment procedure
PSUR	Periodic safety update report



Acronym	Name
R&D VS	Research & Development Value Stream
RPM	Regulatory procedure management
RW Metadata	Real-world metadata
SA	Scientific advice
SAM	Serverless application model
SAFe	Scaled Agile Framework
SIAMED	An information system for the management of regulatory procedure for centrally authorised products
SME	Micro, small and medium-sized enterprises
SME	Subject matter expert
SMS	Substance Management Service
SOC	Security Operations Centre
SPOR	Substance, Product, Organisation and Referential
SSA	Signal and safety analytics
TRIP	$\underline{I}$ opics, $\underline{R}$ elationships, $\underline{I}$ mpact assessment, $\underline{P}$ roposal generation - for collaborative horizon scanning and work on regulatory science
TLM VS	Technology Lifecycle Management & Information Security Value Stream
UPD	Union Product Database (for veterinary products)
UPhV	Union Pharmacovigilance Database (for veterinary products, formerly known as EVVet3)
Value Stream	Value streams represent the series of steps that an organization uses to implement solutions that provide a continuous flow of value to the business/customer
VSM	EMA Value Stream Manager (VSM) is a "Servant Leader and Coach" for the Value Stream teams
VSO	EMA Value Stream Owner (VSO) has primary responsibility for business outcomes, including delivery of business outcomes, in their value stream
XEVMPD	eXtended EudraVigilance Medicinal Product Dictionary  EMA