Rapid Alert system for human Tissues and Cells (RATC) and for human Blood and Blood Components (RAB)

Summary of 2020 activities
Introduction

The rapid alert platforms for blood (RAB) and for tissues and cells (RATC) give Member States' competent authorities the possibility to create and launch alerts to each other and/or to request information in case of an alert or crisis involving more than one Member State. The systems facilitate the communication of information needed to allow competent authorities in other Member States to rapidly assess risks and take adequate and timely measures.

DG SANTE hosts these two platforms, maintains the standard operating procedures (SOPs) and manages users from the national competent authorities. These national users are the ones who draft, launch and close the alerts.

This report provides an overview of the functioning of both systems and alerts submitted in 2020.

Background

Article 8 of Directive 2006/86/EC\(^1\) requires the Member States' competent authorities for human tissues and cells to "communicate to each other and to the Commission, such information as is appropriate with regard to serious adverse reactions and events, in order to guarantee that adequate actions are taken."

Article 9 of Directive 2005/61/EC\(^2\) regarding communication of information between Member States' blood competent authorities and to the Commission requires that Member States "ensure that their competent authorities communicate to each other such information as is appropriate with regard to serious adverse reactions and events in order to guarantee that blood or blood components known or suspected to be defective are withdrawn from use and discarded."

The rapid alert platform for human tissues and cells (RATC) was initiated in 2013 and the rapid alert platform for human blood and blood components (RAB) was initiated in 2014, in order to provide the Member States' competent authorities and the European Commission with an effective and secure tool for the exchange of information for situations in which there is a suspicion of serious health risks associated with tissues, cells, blood and blood components distributed across borders.

The system has been used in parallel with existing national vigilance systems, which collect and manage alerts on human tissues, cells, blood and blood components donated and used within a Member State. Additionally, messages can be communicated regarding problems in related sectors (e.g. medical devices, human or veterinary medicinal products, human organs intended for transplantation) which might imply a risk for the quality and safety of blood, tissues or cells.


**RATC alerts**

The criteria established by the Member States and the European Commission for encoding rapid alerts in the RATC system remained unchanged in the reporting period (e.g. the need for immediate/urgent consideration or follow-up measures in two or more Member States; known or potential risk to patients; issues of a serious or potentially serious nature; potential public health risk to other countries).

Four types of rapid alert were defined and used as follows:

1) **Quality and Safety Defects** are understood as alerts requiring field corrective actions (e.g. recall, quarantine, discard, etc.) of the concerned human tissues/cells potentially affecting patient safety in other Member States.

2) **Information Notices** are defined as alerts related to corrective actions issued in the medical device sector, medicinal products sector or other sector(s), which were of relevance to the tissues and cells sector.

3) **Illegal and fraudulent activities** are defined as alerts used to notify Member States and the European Commission of the possible presence in the distribution network of tissues or cells resulting from actual or suspected illegal and fraudulent activities in the procurement, testing, processing, packaging, distribution, labelling, import/export or promotion of human tissues or cells.

4) **Epidemiological Notices** are alerts related to the development of significant epidemiological situations (e.g. disease outbreaks) which may have cross-border implications in the field of tissues and cells intended for human application.

**Bilateral inquiries** are defined as rapid ways of communication between competent authorities of only two Member States related to any type of alert to be used in particular situations:

- the need to substantiate/confirm information related to a potential rapid alert before the official submission in the RATC system;

- any other situation which is deemed appropriate for such an alert.

At a later stage, an inquiry can be either closed or converted into another type of alert.

The RATC Standard Operating Procedures (SOP) provide guidance on when and how Member States' competent authorities should inform each other.

**Rapid alerts reported in RATC during 2020**

In the interest of openness and transparency to regulatory authorities, professional organisations and other interested parties, the communications via RATC system, reported by the Competent Authorities, are presented below.

A total of 30 alerts were launched in 2020: 25 alerts were encoded in relation to quality and safety defects of tissues and cells (all from DK), 2 alerts were encoded as epidemiological notice (AT 1, NL 1), 2 alerts were encoded as information notice (ES 1, IT 1) and one as “other” (NL). There were no bilateral enquiries. There was a significant
reduction in the number of submitted alerts compared to previous years, almost certainly
due to the fact that activities in tissue establishments were dramatically reduced during
the lockdowns imposed by national governments during the Covid-19 pandemic.

All the alerts encoded as quality and safety defects concerned sperm donations identified
as posing a risk for transmission of genetic disease. Authorities limited further
distribution and use of the donations concerned.

The epidemiological notices encoded concerned: 1) a West Nile Virus outbreak in the
Netherlands and 2) the implementation of preventive measures against West Nile Virus
transmission (donor surveillance and/or deferral and donation testing) in Austria. The last
one was also reported to the RAB network.

Spain also reported an outbreak of West Nile Virus, but as an information notice. The
alert detailed the safety recommendations issued regarding organs, tissues and cells for
clinical use, i.e. donor surveillance and/or deferral and donation testing.

The Netherlands, on behalf of the World Marrow Donors Association (WMDA), reported
in the “other” category delays in the transport of hematopoietic stem cells due to travel
restrictions imposed as measures against Covid-19. WMDA strongly recommended
thorough planning of the transport (including cryopreservation on delivery) and increased
coordination between collection and transplant centres.

Italy also reported difficulties linked to international transport of hematopoietic stem
cells units in the information notice category.

DG SANTE and DG HOME responded to these issues, ensuring that the green lanes
established at the borders to allow free circulation of critical goods were also made
available as channels to facilitate the delivery of substances of human origin within the
EU.

**RAB alerts**

The RAB Standard Operating Procedures (SOP) establish the criteria for encoding rapid
alerts in the RAB and provide guidance on when and how Member States should
communicate with each other. These have been defined by the Member States and the
European Commission. They concern the need for immediate/urgent consideration or
follow-up measures in two or more Member States, a known or potential risk to patients,
issues of a serious or potentially serious nature and potential public health risk to other
countries.

Three types of rapid alert were defined and used as follows:

1) **Quality and Safety Defects** are understood as alerts requiring field corrective actions
(e.g. recall, quarantine, discard, etc.) for the blood or blood components that might affect
patient safety in other Member States.

2) **Information Notices** are defined as alerts related to field corrective actions performed
in the medical device sector, medicinal products sector or other sector(s), which are of
relevance to the blood and blood components sector.

3) **Epidemiological Notices** are alerts related to important epidemiological developments
(e.g. disease outbreaks) which may have cross-border implications in the field of blood
donation and transfusion.
A fourth type of alert, a bilateral communication, is also possible. Bilateral inquiries are defined as rapid ways of communicating between competent authorities of only two Member States related to any type of alert to be used in particular situations:

- the need to substantiate/confirm information related to a potential rapid alert before the official submission in the RAB system;
- any other situation that is deemed appropriate for such an alert.

At a later stage, an inquiry can be either closed or converted into another type of alert.

**Rapid alerts reported in RAB during 2020**

In the interest of openness and transparency to regulatory authorities, professional organisations and other interested parties, the communications via the RAB system, reported by the competent authorities, are presented below.

A total of 11 rapid alerts were encoded in RAB, 8 related to epidemiological notices, one to quality and safety issues and two were information notices. These were issued by the following seven Member States: AT(1), BG (1), ES (3), IT(3), RO(1), SK(1) and NL (1).

In analogy with the RATC, the number of alerts submitted to the blood network almost halved compared to 2019.

Six of the epidemiological notices encoded concerned the implementation of preventive measures against West Nile Virus transmission (donor surveillance and/or deferral and donation testing) in Austria, Netherlands, Romania and Italy. Austria reported the same alert to the RATC network. Italy encoded two more epidemiological notices regarding preventive measures against Dengue and Covid-19 transmission (donor surveillance and/or deferral and donation testing).

Two alerts were encoded as an Information Notice (from Spain and Slovakia) addressing donor selection, preventive measures in blood collections, sufficiency in blood products, additional safety measures and haemovigilance during the Covid-19 pandemic. Belgium encoded one alert as a Quality and Safety issue concerning an increased risk for false negative results when testing HIV antibodies on low-reacting samples after changing detergents in the kit solutions. This test is used as a second-line test after initial screening. Investigation was started and triggered follow up actions.

**Conclusions**

In comparison with 2019, the number of alerts has considerably decreased for both tissues & cells and blood & blood components.

The activities of the Member States in the rapid alert platforms, RAB and RATC, have focused on blood, tissues and cells that are distributed between Member States in Europe and on exchanges of information and description of urgent measures to be taken. While most of the alerts for tissues and cells concerned quality and safety defects, epidemiological notices were the main category of alert in the blood sector.
Once more, the platforms proved to be an effective tool to respond to the needs of authorities for communication and information dissemination in relation to immediate health threats.