1. Introduction

Regulation (EC) No. 1223/2009 on cosmetic products (‘Cosmetics Regulation’)¹ created the basis for a uniform approach to the management of serious undesirable effects (SUEs) attributable to the use of cosmetics. It provides for notification of SUEs without delay to the Competent Authorities of the Member State where the effect in question occurred, as well as the notification of any corrective measures taken by the Responsible Person or Distributor. Data on SUE become part of the Cosmetics Product Safety Report (CPSR)² and have to be made available to the public³.

In order to facilitate the implementation of Article 23 of the Cosmetics Regulation, which constitutes an essential part of a cosmetovigilance⁴ system, and to establish a management and communication system on SUE throughout the EU, the Commission, in conjunction with Member States and industry, established the following guidelines describing the system. Their aim is to ensure harmonized notification of SUE by the Responsible Person or Distributor and follow-up on SUE notifications by Competent Authorities, Responsible Persons or Distributors.

2. Notification and transmission of SUEs

2.1 Definitions

The Cosmetics Regulation defines undesirable effects as “adverse reactions for human health attributable to the normal or reasonably foreseeable use of a cosmetic product”.⁵

Serious undesirable effects are defined as “undesirable effects which result in temporary or permanent functional incapacity, disability, hospitalisation, congenital anomalies or an immediate vital risk or death”.⁶

Taking into account the definition of an SUE, the word "serious" is not synonymous with severe. "Severe" is used to describe the intensity (severity) of the effect as in mild, moderate or severe. Seriousness is used to describe the patient/event outcome or action.

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¹ OJ L 342, 22.12.2009, p. 59
² Annex I of Regulation (EC) No. 1223/2009 on cosmetic products
³ Article 21 of Regulation (EC) No. 1223/2009 on cosmetic products
⁴ Cosmetovigilance is defined by the collection, evaluation and monitoring of spontaneous reports of undesirable events observed during or after normal or reasonably foreseeable use of a cosmetic product. Together with other tools, cosmetovigilance contributes to post market surveillance.
⁵ Article 21. (o) of Regulation 1223/2009
⁶ Article 21. (p) of Regulation 1223/2009
As a consequence, before the notification or the transmission of the SUE, Responsible Persons, Distributors and Competent Authorities must ensure that the undesirable effect meets the seriousness criteria.

2.2 Causality assessment

Causality assessment is an analysis of causal association, on a case-by-case basis, in an attempt to determine the probability that a serious undesirable event is attributable to a well identified product used by an end user.

The causality assessment method described in the Annex 1 to this guideline provides a state-of-the-art approach to determine whether a notified serious undesirable event is considered to be attributable to the use of a cosmetic product.

The causality assessment relates to the effect on an individual end user; it does not provide any evaluation of the risk of a product to the general population. The likelihood of causality should be obtained from the use of a standardized method for causality assessment (see Annex 1).

The aim of this method is to provide a basis for a common understanding and uniform approach to the performance of causality assessments for serious undesirable events to cosmetic products.

In order to perform the causality assessment, information is needed on the serious undesirable event and on the product. To this end, an exchange of all the relevant information between the Responsible Person, the Distributor and the Competent Authority is crucial.

Notifications by the Responsible Person should include a causality assessment, which should be reviewed by the Competent Authority.

Notifications by the Distributor should, if possible, include a causality assessment, which should be reviewed by the Competent Authority. In any event, the Distributor should gather all the available information on the case in order to allow the Responsible Person and/or Competent Authority to make the causality assessment.

Causality assessments for cases reported directly to Competent Authorities should be made preferably by the Authorities. If this is not possible, the Authorities should inform the Responsible Person and exchange all available information to allow a causality assessment to be performed by the Responsible Person without delay.

The person responsible for the causality assessment should be someone who is experienced in complaint handling and who has an appropriate professional background. In certain cases it may be advisable to seek the support of an external or in-house healthcare professional in making the causality assessment in order to obtain a high degree of confidence in the result.

It is possible that the outcome of an initial assessment may change at a later stage in the process as a result of additional information obtained from detailed

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7 Event is a noxious and unintended reaction that occurs in humans using or exposed to a cosmetic product without prejudging in advance of a link between a cause and an effect.
questionnaires or from medical investigation. A causality assessment should only be considered “final” if it is unlikely that further information will be obtained which might change the assessment.

2.3 Scope of notification of SUEs

The Cosmetics Regulation requires the notification by the Responsible Persons and Distributors of all serious undesirable effects which are known to them or which may be reasonably expected to be known to them.

Due to their potential medical seriousness, all SUE cases, except those classified as ‘excluded’ in causality assessment within the timeframe indicated in section 2.4.3, should be notified and the information on these cases should be kept available by the Responsible Person for the Competent Authority of the Member State in which the Responsible Person is established.

The act of notifying a SUE to a Competent Authority is not to be construed as an admission by the company of liability for the SUE and its consequences.

2.4 Requirements for notification and transmission of SUEs

2.4.1 Notification /Transmission Forms

Three different forms were drawn up, enabling a structured and harmonised submission of all important factors related to the SUE, as well as relevant ancillary information (report reference number, outcome of causality assessment, status of notification: initial vs. follow-up, etc.)

The use of the following forms (see Annex 2) is strongly encouraged:

- SUE Form A: Responsible Persons or Distributors notifying SUEs to the Competent Authorities;
- SUE Form B: This form is completed by the Competent Authority and attached to SUE Form A to provide a brief summary and perspective of the case when the Competent Authority transmits SUE Form A to other Competent Authorities and to the Responsible Person. The transmission to the Responsible Person is mandatory when the initial notification comes from a Distributor and it is highly recommended in follow-up and final transmissions when the initial notification comes from the Responsible Person; and
- SUE Form C: Competent Authorities transmitting SUEs reported by health professionals or end users to other Competent Authorities and the Responsible Person.
Flowcharts for notification scenarios

1. **SUE initially received by the Responsible Person or the Distributor**

   SUE Form A: Responsible Persons or Distributors notifying SUEs to the Competent Authorities;
   SUE Form B: Transmission Form for National Competent Authority (accompanying Form A to provide a brief summary and perspective of the case, when transmitting information to other EU Competent Authorities and the Responsible Person):
   - to be sent to other EU Competent Authorities when received initially by the Responsible Person or the Distributor (recommended to be also sent to the Responsible Person)
   - to be sent to the Responsible Person when the initial notification comes from the Distributor

2. **SUE initially received by a National Competent Authority**

   SUE Form C: Competent Authorities transmitting SUEs reported by health professionals or end users to other Competent Authorities and the Responsible Person
The forms are designed not only for the initial notification or transmission, but also for follow-up and final conclusions. Not all the information listed in the forms may be available at the time of the initial notification. However, the initial notification should be carried out if the following minimum information is available:

a) an identifiable reporter;

b) the nature of the alleged SUE and the date of its onset; and

c) the name of the cosmetic product concerned enabling its specific identification.

If the minimum information cannot be obtained, the notifier should continue to undertake all reasonable efforts to obtain the information and notify without delay as it becomes available. The existence of SUE cannot be confirmed unless a minimum amount of information can be obtained.

The list of Competent Authorities will be compiled and made available to the public by the European Commission.

2.4.2 Identification /traceability of SUEs

Each Member State and the Responsible Person or Distributor should be able to unambiguously identify the cases which are forwarded to them.

A common European identification system should be used by Competent Authorities for their management of cases of SUEs when they first receive them (e.g. OECD coding for the country of origin, the year of reporting and the serial number of the concerned case). To avoid duplication, and to manage the follow up information of SUE appropriately, both the Company and Competent Authority Case Identification Numbers should be printed on the documents exchanged on these cases.

2.4.3 Timeframes

For the interpretation of the delays referred to in points 1 (without delay) to 4 (immediately) of Article 23 of the Cosmetics Regulation, the timeframes should be understood as being within 20 calendar days from the date on which any employee of the company or of the Competent Authority, whatever their role or function, becomes aware of the SUE.

2.5 Principles of interaction between the Responsible Person, Distributor and Competent Authorities

The Cosmetics Regulation makes provision for an exchange of information between the Member States’ Competent Authorities and the company (Responsible Person or Distributor) whose product is concerned by the SUE notification.

The Responsible Person or Distributor should exchange all available information that is relevant to the assessment of the case. Additional information deemed necessary by the Competent Authority should be provided on request.
Prior to forwarding information to other Competent Authorities, the Competent Authority receiving a SUE notification should verify whether the case fulfills the seriousness criteria described in Chapter 2.1 and whether the required minimum level of information is available (Chapter 2.4.1).

Where several products are suspected, the Competent Authorities should involve all Responsible Persons concerned.

To ensure the efficiency of the system and to avoid duplication, it is recommended that the Responsible Person receives a copy of the transmission form disseminated to the other Competent Authorities. If other significant information relevant to the case, including its final conclusion, is exchanged between Competent Authorities, it is also recommended that the Responsible Person should be informed.

In particular, the Responsible Person should have the opportunity to review and comment on the causality assessment. If there is no consensus between the Competent Authority and the Responsible Person on the causality assessment, this disagreement should be noted in the transmission of the SUE to the other Competent Authorities.

It is recommended that any communication to the Responsible Person or between Competent Authorities on a notified SUE should be channelled through the Competent Authority which originally received the notification.

Distributors are under a legal obligation to notify to the authorities any SUEs that are reported to them. It is acknowledged that they may not have the same level of information on the product as is available to the Responsible Person and they may find it difficult to provide the full information expected in a SUE notification. The Distributor can inform the Responsible Person in order to collaborate on the SUE notification, provided that the timeframes referred to in Chapter 2.4.3 are respected.

3. Transmission of information on SUEs between Competent Authorities

3.1 Principles

The scope and objective of information exchange / transmission on SUEs between authorities is to facilitate post-market surveillance in order to ensure that the provisions of the Cosmetics Regulation are respected.

Prior to their transmission to all Competent Authorities, the causality of SUEs should be determined by means of the common method referred to in Chapter 2.2.

Likewise, any changes in the outcome of the causality assessment, based on relevant follow-up information to a case, should be transmitted to the Competent Authorities, including assessments which ultimately rule out a link between the product and the SUE.

3.2 Information Exchange Network among Competent Authorities

The information exchange concerning SUEs among Competent Authorities of the Member States will be carried out via the Communication and Information Resource Centre for Administrations, Businesses and Citizens (CIRCABC).
3.3 Data privacy protection and confidentiality issues

All persons involved in SUE notification and transmission should be familiar with and discharge obligations with regard to the collection, use and disclosure of personal information in accordance with the national regulations transposing the EU Personal Data Protection Directive\(^9\). In particular, end users and / or notifiers (e.g. health professionals) should not be identified by their name or address when notifying a SUE or when disseminating a notification among Competent Authorities.

All communications concerning SUEs between Responsible Persons and Competent Authorities, between Distributor and Competent Authorities, between Responsible Person and Distributor, or between different Competent Authorities should guarantee the confidentiality of the information. The reception and the storage of the SUE forms received should be accessible to clearly identified authorized persons only, in accordance with internal Standard Operating Procedures.

4. Subsequent actions

The main purpose of subsequent actions is to maintain the protection of health and safety of cosmetics users by reducing the likelihood of recurrence of an SUE. This includes, where appropriate, corrective measures and the dissemination of information which could be used to prevent such repetitions and which should be proportional to the nature and/or frequency of the SUE.

It should be stated that notification of a SUE does not necessarily indicate a serious risk or non-compliance of the product.

Besides the evaluation of isolated cases, ideally the validation of a signal\(^{10}\) and the measure of its impact should be performed. This requires further investigations using other sources of information, the identification of possible risk factors and the characteristics of the population exposed.

Therefore, care should be taken when evaluating spontaneous reports\(^{11}\), especially if a comparison is made between different countries or companies. The data accompanying spontaneous reports and the rate at which cases are reported is dependent on many factors. In order to minimize bias, a specific analysis and evaluation of medically validated SUEs should be considered and compared with non-medically validated cases.

4.1 Subsequent actions by a Responsible Person

4.1.1 Analysis of the data

A human health issue could be identified from one report or, more likely, from several similar SUE reports associated with the same product. Where necessary, a trend analysis that takes into consideration the nature, severity and/or frequency should be

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\(^9\) OJ L 281, 23.11.1995, p. 31

\(^{10}\) A signal may be defined by the onset of an unexpected modification of a pre-existing level of reporting rates, including qualitative or quantitative modifications.

\(^{11}\) In vigilance systems spontaneous report refers to unsolicited communication by end users or by health care professionals to a company, regulatory authority or other organization that describes one or more suspected health related events in a person who has used one or more cosmetic products.
performed. Other factors could include possible predisposing factors on the part of the end users who had experienced the undesirable effect.

When a human health issue is thereby identified, further analyses should be carried out to establish, where possible, the potential mechanism of the undesirable effect.

4.1.2 Inclusion in the Cosmetic Product Safety Report

Annex I of Cosmetics Regulation requires inclusion in the Cosmetic Product Safety Report of “All available data on the undesirable effects and serious undesirable effects to the cosmetic product or, where relevant, other cosmetic products. This includes statistical data.”

Specific guidance on this matter is given in separate EC guidelines (ref: Annex I Guidelines)

4.1.3 Information to the public

The Cosmetics Regulation requires that existing data on undesirable effects and serious undesirable effects resulting from the use of the cosmetic product are made accessible to the public by any appropriate means\(^\text{12}\).

Although that information has to be made accessible to the public on request, it does not have to be published. The contents should be presented in a consistent fashion and follow the recommendations described in the Guidelines for Information to the Public (reference to be updated).

Any communication on cosmetovigilance data should take into account the understanding level of the readers. Data should be provided on causality levels and levels of seriousness. To be meaningful, cosmetovigilance data should not be presented in isolation, but should be put into perspective with market data.

4.1.4 Corrective action

Where necessary, a number of actions may be undertaken by a company following assessment of the post marketing surveillance data, together with other sources of safety data. The measures taken should be proportional to the nature and/or frequency of the SUE and be subject to the same rigorous risk assessment exercise conducted by Competent Authorities (see below). These measures may include a change in usage instructions, labelling, warnings, changes to the formula, recall or withdrawal of the product, or any further action necessary to protect the health of the end user. If a SUE requires corrective measures, these have to be notified to the same Competent Authority to whom the SUE had originally been notified. This Competent Authority has to inform other Competent Authorities in the Union.

4.2 Subsequent actions by Competent Authorities

Actions subsequent to SUE notifications can be taken by Competent Authorities for the purposes of in-market surveillance, market analysis, evaluation, and end user information in the context of Articles 25, 26 and 27 (non-compliance and safeguard clause).

\(^\text{12}\) Article 21 of Regulation (EC) No. 1223/2009 on cosmetic products
4.2.1 Evaluation of trend or signal detection

Identification by a Competent Authority of a signal or a trend based on the report of SUEs could lead to a specific enquiry in the country concerned; the Responsible Person should be informed of the enquiry so that they can provide the investigating Competent Authority with the information needed to evaluate the trend or signal. The analysis of the signal should follow state-of-the-art risk assessment principles, e.g. those described by the International Risk Governance Council\textsuperscript{13}.

If Competent Authorities decide to investigate further at European level, the Responsible Person and the European Commission should be informed.

Except in the case where immediate action is necessary on the grounds of a serious risk to human health, the Responsible Person should be given the opportunity to put forward his viewpoint before any decision is taken.

4.2.2. End user information by Competent Authorities

Periodic bulletins on post-marketing surveillance data from cosmetics may be issued by Competent Authorities, particularly on their respective websites. If data on SUEs including the outcome of causality assessments and statistical analysis are published through this medium, the Responsible Persons of the companies concerned should be duly informed ahead of such publication if the commercial name of the product is mentioned.

The risks of a dissemination of isolated cases of SUEs to the public should be carefully examined. Any communication on cosmetovigilance data should take into account the level of comprehension of the readers. In order to be meaningful, cosmetovigilance data should not be presented in isolation, but should be put into proper perspective. Data should be provided on causality levels and degree of seriousness.

Accurate and timely communication of emerging data on risk is an essential part of cosmetovigilance. Risk communication is an important stage in risk management as well as a risk minimisation activity. End users and healthcare professionals need accurate and effectively communicated information about the risks associated with the cosmetic products and other factors influencing these risks. Because of the importance of risk communication, it is recommended that appropriate experts should be consulted.

Annex 1:

CAUSALITY ASSESSMENT OF UNDESIRABLE EFFECTS CAUSED BY COSMETIC PRODUCTS
SUMMARY

A causality assessment method for undesirable effects potentially caused by cosmetic products was developed by a group of experts.

The aim of this reproducible, rational, harmonised and standardised method is to assess cause-and-effect relationships between cosmetic products and given clinical and/or paraclinical effects.

The method is based on six criteria, divided into two groups, which are used to calculate a chronological score and a semiological score.

As a rule, the method must be used separately for each cosmetic product, without taking into account the level of causality of the associated products.

The level of causality is determined using a decision table in which the scores are combined.

The method offers five levels of causality assessment: very likely, likely, not clearly attributable, unlikely and excluded.
Health vigilance systems have two fundamental objectives:

- to record and identify undesirable effects for humans, directly or indirectly caused by a technique, treatment or product;
- to analyse the data collected in order to put in place corrective or preventative measures.

The vigilance process may serve various activities in various fields: improving knowledge, epidemiology, surveillance, signal detection and alerts.

Undesirable effects can occur at random or be linked to specific circumstances or combinations of circumstances or to specific characteristics of each individual.

For a number of reasons, particularly on epidemiological grounds, it can be useful to list already known effects in order to determine their frequency and analyse thoroughly their determinants. By combining their frequency and severity, it is possible to determine the criticality of the undesirable effects, which is one of the central factors in risk management.

However, it is essential to be able to identify undesirable effects irrespective of current scientific knowledge, particularly the scientific knowledge of the reporter and monitor. It is therefore vital not to reject reported undesirable effects on the ground that no causal link can be established.

All healthcare professionals accept that undesirable effects caused by health products cannot be assessed in a purely subjective fashion. The related consequences in terms of health and industrial decisions are significant enough to justify use of an objective and specific diagnostic method.

The aim of such 'causality assessment methods' is to estimate the extent of the cause-and-effect relationship between one (or more) health product(s) and the occurrence of an undesirable effect.

As this is a standardised approach, its main advantage is to eliminate any differences of opinion between individual observers [1 to 3]. Such methods are widely applied for most health products in France and are recommended at European level for cosmetic products (Colipa [7], Council of Europe [8]). In France, the first causality assessment method to be used and published was the pharmacovigilance causality method [4, 5], but there was no harmonised French method for cosmetic products.

At the request of AFSSAPS (the French Health Products Safety Agency), a group of experts developed a causality assessment method suited to the specifics of the undesirable effects attributable to use of cosmetic products.

The approach applied to develop this tool established a number of principles:

- Objective: to develop a generic method, applicable to all cosmetic products and all kinds of observed effects.
- Aims of the method: to allow rating of the level of relationship between a suspected cosmetic product and an observed undesirable effect.
- Identification of relevant criteria to establish a cause-and-effect relationship.
• Analysis of these criteria based on the expected outcomes and the weightings to apply to them.
• Combination of these criteria using a decision table.
• Dual validation of the method:
  - theoretical, by checking the relevance of the outcomes obtained;
  - experimental, by using the method in real-life situations.

As is the case for all causality assessment methods [6], implementation of this method:
• is possible only once a minimum amount of information has been collected;
• must be conducted independently for each cosmetic product used before occurrence of the undesirable effect;
• might require specialist medical assessment — this is recommended in complex cases, or if the impact on the user's health is deemed serious.

This search for information should make it possible to identify any other cause, which is a more likely origin of the undesirable effect than the cosmetic product.

The group of experts established a set of intrinsic criteria, involving no data other than those collected on the individual case, for calculating two types of scores:
• a chronological score; and
• a semiological score.

**Chronological score**

The chronological score is calculated from the information on the time sequence between use of the cosmetic product and occurrence of the symptoms.

The time sequence between use of the cosmetic product and occurrence of the alleged undesirable effect may be:
• compatible, i.e. usual given the reported symptoms;
• only partially compatible, i.e. unusual given the reported symptoms;
• unknown;
• incompatible, whenever the clinical or paraclinical effect occurred before the cosmetic product was used or whenever the period before the observed symptoms appeared is too short.
  If the time sequence is inconsistent, the undesirable effect cannot be attributed to use of the cosmetic product.

**Semiological score**

The semiological score is calculated from the information on the nature of the undesirable effect and on the results of any specific additional examinations that were performed or of re-exposure to the cosmetic product.
a) Symptomatology

Symptomatology is defined as a set of symptoms, recorded as exhaustively as possible during the case investigation, enabling a diagnosis to be put forward. Absence of diagnosis does not prevent use of this method.

It points to use of a cosmetic product whenever the symptoms observed are appropriate to the nature of the product or to its method of use in terms of location, effect or evolution.

It is otherwise only partially or not at all evocative.

In certain cases, factors that might have contributed to the undesirable effect, i.e. to attenuating or accentuating its clinical expression, may come to light when this information is collected. Although these factors may play a significant role, for the sake of simplification they have not been taken into account in this method.

b) Additional examinations (AE)

Any additional examinations must be reliable and specific to the observed effect and must be performed by specialist physicians.

The results of these examinations are rated as follows:

- AE (+): positive;
- AE (-): negative;
- AE (?): if no examinations were performed or if the results were ambiguous.

c) Re-exposure to the cosmetic product (R)

After the decurrence of clinical signs, there are three possibilities if the effects recur after re-exposure to the cosmetic product, whether accidental or not:

- R (+, positive): the initial symptomatology recurs with the same intensity or with a higher intensity when the user is re-exposed to the product;
- R (?): there is no re-exposure to the product or the conditions of re-exposure are not identical to those of the initial exposure;
- R (-, negative): the effect does not recur when the user is re-exposed to the product.

For re-exposure to be considered negative, it must occur under similar conditions of use of the cosmetic product (identical product, identical procedure, identical duration, etc.) without causing an identical undesirable effect (identical symptoms and location, identical time sequence before occurrence, etc.).

These scores, combined in a decision table (Table 1) or a decision tree (Table 2), produce five levels of causality: very likely, likely, not clearly attributable, unlikely and excluded.
In this decision table, in principle causality is ‘excluded’ if the time sequence before the effect appears is considered incompatible.

When other aetiologies might account for an undesirable effect observed, these weaken the alleged link between the cosmetic product and the undesirable effect in question and, consequently, the causality is downgraded by one level, but never ‘excluded’.

In any case where another aetiology explaining the undesirable effect observed is demonstrated, medically validated and documented, the alleged link between the relevant cosmetic product and the undesirable effect in question is excluded in this particular case. This other aetiology must be medically validated by a physician specialising in the relevant organ and, whenever possible, be reported in writing. The excluded cases will be regularly re-assessed as scientific knowledge progresses.

**Table 1: Decision table**

<table>
<thead>
<tr>
<th>Symptoms</th>
<th>EVOCATIVE of use of the cosmetic product</th>
<th>ONLY PARTIALLY OR NOT EVOCATIVE of use of the cosmetic product</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>R and/or AE +</td>
<td>R and/or AE -</td>
</tr>
<tr>
<td><strong>Time sequence between exposure and occurrence of the symptoms</strong></td>
<td>R and/or AE ?</td>
<td>R and/or AE +</td>
</tr>
<tr>
<td>Compatible</td>
<td>Very likely</td>
<td>Likely</td>
</tr>
<tr>
<td></td>
<td>Likely</td>
<td>Not clearly attributable</td>
</tr>
<tr>
<td>Only partially compatible or Unknown</td>
<td>Likely</td>
<td>Unlikely</td>
</tr>
<tr>
<td></td>
<td>Not clearly attributable</td>
<td>Not clearly attributable</td>
</tr>
<tr>
<td>Incompatible</td>
<td>Excluded</td>
<td>Excluded</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Excluded</td>
</tr>
</tbody>
</table>
Table 2: Decision tree

Symptoms

- **Chronology compatible**
  - Additional examinations and/or re-exposure negative
  - Additional examinations and/or re-exposure not performed or equivocal results
  - Additional examinations and/or re-exposure positive
  - Additional examinations and/or re-exposure not performed or equivocal results

- **Chronology only partially compatible or unknown**
  - Additional examinations and/or re-exposure not performed or equivocal results
  - Additional examinations and/or re-exposure negative

**Not clearly attributable**: If the symptoms are not evocative (not suggestive of the product effect), the final level of causal relationship is decreased by one degree (very likely to likely, likely to not clearly attributable, not clearly attributable to unlikely).

**Compatible chronology**: A time sequence between product use and the occurrence of symptoms as well as between stopping product use and clearing up of the symptoms which is plausible from a medical viewpoint and can be reasonably anticipated for this kind of product use and undesirable event. If the chronology is not compatible the causal relationship is **excluded**.
This decision table was used to establish the following definitions:

<table>
<thead>
<tr>
<th>Causality</th>
<th>Criteria</th>
</tr>
</thead>
</table>
| **Very Likely**    | - the clinical symptoms evoke use of the product;  
                      - the time sequence between use of the product and occurrence of the symptoms is compatible;  
                      - and the specific additional examinations performed are positive and relevant\(^1\) or the re-exposure to the product is positive\(^2\). |
| **Likely**         | - the clinical symptoms evoke use of the product;  
                      - the time sequence between use of the product and occurrence of the symptoms is compatible;  
                      - and there are neither any relevant specific additional examinations\(^1\) nor re-exposure\(^2\) or otherwise the results of re-exposure or the results of the specific additional examinations performed are ambiguous.  
                      Or:  
                      - the clinical symptoms evoke use of the product;  
                      - the time sequence between use of the product and occurrence of the symptoms is only partially compatible or unknown;  
                      - and the specific additional examinations performed are positive and relevant\(^1\) or the re-exposure to the product is positive\(^2\).  
                      Or:  
                      - the clinical symptoms only partially evoke or do not evoke use of the product;  
                      - the time sequence between use of the product and occurrence of the symptoms is compatible;  
                      - and the specific additional examinations performed are positive and relevant\(^1\) or the re-exposure to the product is positive\(^2\). |
| **Not Clearly Attributable** | - the clinical symptoms evoke use of the product;  
                      - the time sequence between use of the product and occurrence of the symptoms is compatible;  
                      - and the relevant specific additional examinations\(^1\) or the re-exposure\(^2\) are negative.  
                      Or:  
                      - the clinical symptoms evoke use of the product;  
                      - the time sequence between use of the product and occurrence of the symptoms is only partially compatible or unknown;  
                      - and there are neither any relevant specific additional examinations\(^1\) nor re-exposure\(^2\) or otherwise the results of re-exposure or the results of the specific additional examinations performed are ambiguous.  
                      Or:  
                      - the clinical symptoms only partially evoke or do not evoke use of the product;  
                      - the time sequence between use of the product and occurrence of the symptoms is compatible;  
                      - and there are neither any relevant\(^1\) specific additional examinations nor re-exposure\(^2\) or otherwise the results of re-exposure or the results of the specific additional examinations performed are ambiguous.  
                      Or:  
                      - the clinical symptoms only partially evoke or do not evoke use of the product;  
                      - the time sequence between use of the product and occurrence of the symptoms is only partially compatible or unknown;  
                      - and the specific additional examinations performed are positive and relevant\(^1\) or the re-exposure to the product was positive\(^2\). |
| **Causality UNLIKELY** | - the clinical symptoms evocate use of the product;  
- the time sequence between use of the product and occurrence of the symptoms is only partially compatible or unknown;  
- and the specific additional examinations(1) or the re-exposure(2) to the product are negative.  
Or:  
- the clinical symptoms only partially evocate or do not evocate use of the product;  
- the time sequence between use of the product and occurrence of the symptoms is compatible;  
- and the specific additional examinations(1) or the re-exposure(2) to the product are negative.  
Or:  
- the clinical symptoms only partially evocate or do not evocate use of the product;  
- the time sequence between use of the product and occurrence of the symptoms is only partially compatible or unknown;  
- and there are neither any relevant specific additional examinations(1) nor re-exposure(2) or otherwise the results of re-exposure or the results of the specific additional examinations performed are ambiguous.  
Or:  
- the clinical symptoms only partially evocate or do not evocate use of the product;  
- the time sequence between use of the product and occurrence of the symptoms is only partially compatible or unknown;  
- and the specific additional examinations(1) or the re-exposure(2) to the product are negative. |
| **Causality EXCLUDED** | - the time sequence between use of the product and appearance of the symptoms is incompatible;  
Or:  
- another aetiology was demonstrated, medically validated and documented. |

(1) The additional examinations performed to objectify an undesirable effect must be specific and relevant: they must follow an established protocol and allow standardised interpretation. These specific and relevant examinations must be clearly defined.

(2) Re-exposure may occur in controlled or uncontrolled fashion. The user may either be spontaneously re-exposed to the product which caused the undesirable effect or otherwise be re-exposed to the product following a specific protocol.
A causality assessment method is a key tool for guaranteeing that a uniform and
rigorous approach is taken for assessing the strength of links between health
products and the occurrence of undesirable effects. This assessment tool is to be
used in conjunction with clinical expertise and knowledge of the relevant products,
which remain essential.

The method proposed here, which is specific to cosmetic products, supplements the
methods commonly used for other health products.

This method must not be considered as definitive per se and must be updated in the
light of the experience gained from large-scale use.

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