Exporting Seafood to the European Union
July 2020 Update
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I. Introduction

A) Scope of the Report:

Given the complexity of the EU legislation, this report provides an overview of key EU legislation governing trade in edible seafood products. It does not intend to answer all questions; additional comments or concerns should be addressed to specific competent authorities (see Points of Contacts at the end of the report).

B) Background:

Twenty-seven countries compose the European Union (EU). The current Member States (MS) are: Austria, Belgium, Bulgaria, Croatia, Denmark, Finland, France, Germany, Greece, Ireland, Italy, Luxembourg, the Netherlands, Portugal, Romania, Spain, Sweden, Latvia, Lithuania, Estonia, Poland, Malta, Cyprus, Hungary, Slovenia, Slovakia and the Czech Republic. The EU population is approximately 508 million people since the accession of Croatia on July 1, 2013.

The European Free Trade Association (EFTA) is the intergovernmental organization of Iceland, Liechtenstein, Norway and Switzerland. These countries although not members of the European Union, fully implement EU food safety legislation.

Following the withdrawal of the United Kingdom from the Union, any reference to Member States shall be understood as including the United Kingdom where Union law remains applicable to and in the United Kingdom until the end of the transition period, according to the Withdrawal Agreement (at least until the end of 2020).

C) The Institutions:

The EU has seven different institutions that function in many ways as the different branches of the US government:

The **European Commission** is the EU executive body. It has three main tasks: to initiate EU policies, to act as the guardian of EU treaties and to supervise implementation of EU law. The Commission is divided in 32 directorates general (DG), of which **DG Mare** and **DG Sante** share responsibility for food safety consumer policy and public health protection. A college of 28 Commissioners who are named by their national governments but should make decisions independently, head the Commission.

The **Council of the EU** consists of Ministers from the National Governments from EU Member States. Each Member State holds the rotating presidency of the Council for six months. The Council and the European Parliament share the responsibility for passing laws and making policy decisions. It also bears the responsibility for what the EU does in the fields of Common Foreign and Security Policy and EU action on several justice and freedom issues. The Council has working parties and permanent or special committees consisting of representatives from Member States. The best known is the Committee of Permanent Representatives of the Member States, or COREPER.
The European Council is the EU institution that defines the general political direction and priorities of the European Union. It consists of the heads of state or government of the member states, together with its President and the President of the Commission.

In addition to the Heads of State, there is a semi-permanent President who serves a two- and half-year term and the rotating President of the Council. While the European Council has no formal legislative power, it is an institution that deals with major issues and any decisions made by it are "a major impetus in defining the general political guidelines of the European Union." The Council meets at least twice every six months.

The European Parliament (EP) is elected every five years by the people of Europe to represent their interests. The core mission of the European Parliament is to pass European laws. It shares this responsibility with the Council of the EU. Proposals for new laws are generated by the European Commission. The EP has the power to dismiss the entire European Commission. The European Parliament has gained authority over time, reaching its peak with the Lisbon Treaty’s “co-decision” authority as well as budget oversight of the European Commission. Co-decision means that no law can be adopted without both European Parliament’s and Council’s consent.

The European Court of Justice rules on disputes involving interpretation and application of the EU treaties and legislation. It makes sure that EU law is interpreted and applied in the same way in all MS. The Court is located in Luxemburg and has one judge from each MS.

The European Central Bank has gained greater prominence during the Euro Zone crisis and the Court of Auditors audits EU finances; its role is to improve EU financial management and report on the use of public funds.

D) What are the different measures?

Regulations:
A Regulation is a law that is binding and directly applicable in all Member States without implementing any new national legislation. Both the Council and the Commission can adopt Regulations.
**Directives:**
A Directive is a law, with specific results to be achieved, that is binding on all Member States. However, each MS can choose how it is to be implemented. In practice, the Commission will issue approved implementing legislation after a Directive is adopted, known as Implementing Measures. Usually, the Commission works with the Member States regarding the details of the implementing measures in to ensure correct implementation of the referred Directive; this an important point, as businesses affected by a Directive must take the national implementing legislation as well as the Directive into account. All Directives include a date by which Member States must transpose the Directive into their national legislation. In case of Member State non-implementation, the Directive remains the legal framework for adjudication. The Commission can act against Member States that have not implemented a Directive on time.

**Decisions:**
A Decision is binding entirely on those to whom it is addressed. No national implementing legislation is required. Both the Council and the Commission can adopt decisions.

**Recommendations:**
A Recommendation has no binding effect -it is not a law. Both the Council and the Commission can adopt recommendations.

**II. How Fishery Policies are handled at the EU Level**

DG Mare is responsible for negotiating international fishing agreements, resources management, aquaculture, fleet management, the Common Fisheries Policy (CFP), ocean governance and maritime affairs. It also sets Autonomous Tariff Quotas (ATQs) that allow third countries such as the U.S. to export fishery products to the European market at a reduced duty. It supports DG Trade, part of which is the EU equivalent to the Office of the US Trade Representative for WTO matters. Some fish species are subject to trade restrictions under the Convention on International Trade of Endangered Species. DG Environment is the “Chef de File” for this subject matter. DG Mare and DG Environment work together closely due to the current status of worldwide fish resources. Fishery products are also subject to measures introduced by DG Agriculture and DG Internal Market and are supervised by DG Sante. DG Agriculture is responsible for the Common Agricultural Policy (CAP) and all “vertical” measures on raw materials. These DGs initiate proposals on all EU measures concerning sanitary legislation and inspection by type of products (beef, pork, poultry, vegetables, seafood, etc.).
DG Grow deals with “horizontal” measures for processed products. Together with DG Sante, they propose legislation on additives, microbiological criteria, colorings, antibiotics, and labeling. All those texts refer to “foodstuffs.” DG Sante oversees all scientific committees that advise DG Grow and DG Agriculture on matters concerning consumer health. DG Sante also includes the Food and Veterinary Office (FVO), which is based in Ireland.

The main responsibilities of the FVO are to monitor the observance of food hygiene, veterinary, and plant health legislation within the European Union, and to help promote confidence in Europe’s food safety to consumers. The FVO is responsible for auditing Member States’ competent authorities and for inspecting third countries’ compliance and/or equivalency to EU legislation.

The European Food Safety Authority (EFSA) was created on January 28, 2002. EFSA covers risk assessment as well as risk communications. The relevant EU institutions maintain risk management responsibility for the EU. Part of EU legislation is defined according to EFSA’s scientific opinions and recommendations.

The primary EU laws that impact US seafood exports are: The Common Fisheries Policy (CFP) and the Food Hygiene Legislation. The CFP establishes a legal framework for the regulation of fisheries and aquaculture activities. It has a direct impact on the EU’s production capacity through fleet and quotas management. Therefore, it can directly affect imports of seafood from third countries such as the US. The Food Hygiene Legislation is the EU’s instrument that guarantees safe food to European consumers. It makes sure that “domestically made” as well as imported food complies with the EU’s minimum hygiene standards.
III. The Common Organization of the Market in Fishery and Aquaculture Products

The Common Organization of the Market in Fishery and Aquaculture Products was first introduced in 1970, reviewed in 1993 and 2000 and last amended in 2013 (Council Regulation 1379/2013). Its purpose is to stabilize the market, to guarantee a steady supply of quality products, to ensure reasonable prices for consumers and support fishermen’s incomes.

The five components of the Common Organization of the Markets are:

- **Marketing standards and consumer information** for fresh products for quality, grades, packaging and labeling for domestic production as well as imports.
- **Producers’ organizations** (voluntary fishermen associations) are officially recognized and are set up to help stabilize markets fluctuations. Their role is to protect fishermen from sudden changes by adjusting supply to demand. They also help to improve product quality and ensure that fishing quotas are respected.
- **Interbranch Organizations and Agreements** are aimed at facilitating a total integration of the sector from producer to consumer.
- **Prices and Intervention** by which certain species cannot be sold below a given price. They can be stored and sold when market improves or processed.
- **Trade with third countries.** In order to ensure an adequate supply of fishery and aquaculture products to the EU intended for the processing industry, the EU adopts regulations providing reduced duty rates, quotas, and autonomous suspensions for specified products.
IV. Exporting Seafood to the EU

A) General Provisions:

As a general principle, seafood is imported into the EU from only approved countries and from approved establishments, e.g., processing plants, factory or freezing vessels, cold storages or brokers. Aquaculture products, including live bivalve mollusks, may be exported from only approved establishments located within approved production zones or areas.

Since 2006, the U.S. Seafood Inspection System has been recognized by the EU as an equivalent of the European Seafood Inspection System. This status does not apply yet to the export of live bivalve mollusks, in whatever form.

This mutual recognition facilitates seafood trade between the U.S. and the EU. Furthermore, it creates a framework under which Member States cannot impose national requirements on U.S. seafood exporters on top of EU harmonized legislation. However, differences of interpretation among Member States can lead to delays at border control posts (BCP).

B) List of Countries:

Commission Implementing Regulation 2019/626 is the list of countries and territories from which imports of fishery products and bivalve mollusks, echinoderms, tunicates and marine gastropods are permitted. One may note that only two U.S. States (Washington and Massachusetts) appear on the list of countries authorized to export bivalve mollusks, echinoderms, tunicates and marine gastropods. This means that, unlike fishery products, the U.S. inspection system for shellfish has not been recognized as totally equivalent to the EU’s inspection system. Similarly, the U.S. only recognizes two EU Member States as being equivalent to the U.S. (The Netherlands and Spain). Negotiations between the U.S. and the EU to resume transatlantic trade in shellfish are currently underway.

However, the exports of wild roe-off scallops to the EU from the U.S. is still authorized, per Article 8 of Regulation 2019/626 referred above. In this case, a regular health certificate for fishery products is required.

C) Approved Establishments

U.S. operators that wish to export seafood to the EU must be approved by and registered with their National competent authority. The Food & Drug Administration (FDA) is the U.S. agency
responsible for the approval of seafood establishments. Once they are approved, U.S. exporters are included on the FDA list, which is updated every quarter. This FDA list is then sent to the EU for validation. **The process can take up to three months.** The list of FDA District Offices in charge of the approval process can be found at: 
https://www.fda.gov/ForIndustry/SmallBusinessAssistance/default.htm

**Although the production is allowed from the date the request for listing a new establishment has been confirmed by the European Commission services, no health certificate can be issued by the competent authority overlooking that establishment before the amended published EU list of approved establishments enters into force.**

**D) Certification**

Since January 1, 2010, each shipment of seafood products must be accompanied by a **Sanitary** **AND a Catch Certificate.** You will find a separate chapter on the catch certificate later in this report.

**Important Notice:** Since June 2009, the U.S. Department of Commerce, NOAA/National Marine Fisheries Service, is the U.S. agency responsible for the certification of fishery and aquaculture products intended for the EU. A health certificate may be issued for goods produced by different establishments but can only be made to one consignee.

A health certificate may be issued for several containers of the same product considered to be a single lot. The health certificate must define the lot. Therefore, a rejection at the point of entry will include all goods covered by the same health certificate, even if only a part of it presents a sanitary or documentary problem. It is acceptable to list fresh and/or live products on the same health certificate. However, frozen products must be listed on a separate health certificate. Health certificates must be issued in one of the official languages of the country of entry into the EU territory, and if necessary, in the language of the country of destination. However, a Member State may consent to the use of one of the 23 official EU languages other than its own.

Since December 14, 2019, the EU has modified the template of official certificates for certain animal and goods. This **Regulation does not** apply to our existing U.S. fishery products certificate (article 4, paragraph 4 of the prementioned Regulation) except for imported products that have not been initially produced/processed in the U.S. and that are re-dispatched to the EU without further processing. It also modifies and restricts the conditions for a replacement certificate; see article 5.
In practice, the Border Control Post (BCP) at the first point of entry into the EU conducts the documentary check and issues a Common Veterinary Entry Document (CVED) in conformity with Commission Regulation 136/2004.

This CVED must be:
1. Either in the language or one of the languages of the border control post where the products are entering the EU, and
2. Either in the language or one of the languages of the destination country.

**Important Notice:**
Since April 1, 2007, Switzerland adopted EU sanitary legislation regarding import requirements for fishery products. Therefore, U.S. seafood shipments must be accompanied by the same health certificate as required by any EU Member State. A health certificate intended for Switzerland may be in French or English.

**E) Import Controls:**
Principles for veterinary checks are laid down in Regulation 2017/625. Inspections of consignments originating from third countries must be carried out on all consignments, at the first point of entry into the EU territory and at approved border control posts.

Import controls are done in three consecutive steps:

1. **Documentary check:** examination of the health certificate;
2. **Identity check:** visual inspection to confirm consistency between documents and products, verification for the presence of required sanitary marks - country of origin, approval number; and,
3. **Physical check:** check of the product itself, organoleptic control, packaging, temperature. This may include sampling and laboratory testing.

Products imported from “harmonized” countries, such as the U.S., are subject to the documentary, identity and physical checks at the approved border control post at the first point of entry into the EU territory. When a consignment satisfies EU requirements, it can be marketed freely in all EU Member States.
While the documentary and the identity checks must be performed on all consignments, the frequency of physical checks is reduced for products from “harmonized” countries. Approximately 20 percent of fish products in hermetically sealed containers, fresh and frozen fish, and dry or/salted products undergo physical checks. For other fishery products, about 50 percent are subject to physical checks. Each import control - one certificate = one control - is subject to inspection fees. In the case of processed food containing animal products, surimi, for example, the European importer must have an “import license” from their customs authorities before the import process begins.

European border control posts may randomly conduct specific analysis on shipments presented to them for clearance. The analyses can target residues, heavy metals or other contaminants. Normally products will not be allowed to leave the BIP pending test results. In cases where samples are taken for analysis and the results will not be known for at least a few days, consignments will not be given a CVED and will not be able to leave the BCP if:

- The testing is for a substance or pathogenic agent which presents a direct or immediate animal or public health risk; or
- The testing has been carried out because of information on previous unfavorable test results.

The veterinary authorities for the area of destination should be advised of any pending test results by TRACES, *Consignments which are sampled as part of the Veterinary Monitoring Directive (VMD) non-statutory sampling plan and those taken as part of routine sampling at the BCP or the national monitoring plan do not need to be detained pending the results of the analysis.*

However, if the tests reveal any contamination, the establishment that sent the shipment in question will be put on “reinforced control status.” This status is then communicated to all Member States as well as to the European Commission through the Rapid Alert System. When an establishment is on reinforced control status, the U.S. establishment’s next ten consecutive shipments (of a commercial size, not sample) to any EU country will be automatically tested. The products will be detained at border control posts until results are received. After ten shipments without positive results, the establishment in question is removed from the reinforced control list. The legal text describing reinforced control procedures is [Commission Implementing Regulation (EU) 2019/1873](https://eur-lex.europa.eu/eli/reg/2019/1873/oj).

If a shipment is refused for non-compliance with EU legislation, the responsible party of the shipment has three options:

1. Destroy the products in question;
2. Re-dispatch these products to a non-EU country; or
3. Return the products to the originating country.

It is important to note that Regulation 882/2004 (Article 21) imposes a number of conditions for the two last options noted above:

1. The new destination has been agreed to by the EU based food business operator, i.e., consignee;
2. The consignee must inform the competent authority of the third country of origin or third country of destination, if different, of the reasons and circumstances that prevented sales of the food within the EU;
3. **And**, when the third country of destination is not the third country of origin, the competent authority of the third country of destination must signal its preparedness to accept the consignment.

**F) Triangular Trade:**

Triangular trade occurs when U.S. products are shipped from the U.S. to other third (non-EU) countries for storage before being re-exported to the European Union later. It also occurs when a product is imported into the U.S for storage or for further processing and then re-dispatched to the EU.

EU sanitary legislation requires:
1) The shipment must be stored in an EU-approved facility in the third country;
2) At the time of re-export to the EU, it must be accompanied by a sanitary certificate from the last country of dispatch, even if the products were not further processed in that country.

This certificate must be based on the info included on the first certificate that was issued by the initial competent authority, if applicable.

If the products are destined to a third country with temporary storage in a specially approved customs warehouse on the EU territory, they need to be presented to the EU border control post of arrival for transit controls. They should at least be accompanied by the health certificate requested by the third country of destination and the BCP will issue a Common Veterinary Entry Document (CVED). A second CVED will be issued when the goods leave the customs warehouse and travel to the country of destination. This is applicable for most fishery products; however, for fish and crustaceans of aquaculture origin, the health certificate needs to be provided to the entry BCP as such transit consignments must fulfill animal health conditions.

Certification procedures for re-dispatch to another third country are then of the responsibility of the member states where the goods are stored. Some may issue a certificate; some will refuse to do so. In any case, please contact NOAA EU office before shipping.

**V. Food and Feed Hygiene Legislation**

Hygiene is part of the European policy on food safety, which also considers other sanitation aspects such as materials in contact with food, labeling, chemical substances, e.g., additives and food colorants, and ionization of foodstuffs, contaminants and residues.

While this Hygiene Package tends to simplify the previous very complex legislation, it also introduces the concept of “responsibility” for the food and feed operators throughout the entire food chain, in other words, “from farm to fork.” This section summarizes the legislation specifically addressing fishery products and bivalve mollusks.
A) Food Hygiene:

The Hygiene Package sets clear and strict rules on the sanitary conditions of foodstuffs, specific sanitation rules for food of animal origin, and specific rules for controls on products of animal origin intended for human consumption. While there are general rules for all food, there are specific measures that apply to fishery products and bivalve mollusks.

Under this updated legislation, imported products will be required to meet the same standards as EU products.

The Hygiene Package is divided into 5 Regulations and Directives:

**Hygiene 1**:

**Hygiene 2**:
European Parliament and Council Regulation 853/2004 are specific sanitary rules for food of animal origin. Specifically, Annex I - definition, and Annex III Section VII & VIII - bivalve mollusks and fishery products. This Regulation has been amended by Regulation 1662/2006. The last amendment modifies the conditions for exports of fishmeal into the EU.

**Hygiene 3**:
Regulation 2017/625 outlines specific rules for the organization of official controls on products of animal origin that are intended for human consumption.

**Hygiene 4**:

**Hygiene 5**:

B) Subsequent Regulations:

U.S. exporters should be aware that Member States may have adopted additional measures that are specific and must be followed in addition to the requirements of the Hygiene Package.

Microbiological Criteria for Foodstuffs:

These criteria are fundamental for a comprehensive food sanitation framework. Regulation 2073/2005 last amended by Regulation 2015/2285. These criteria are applicable to products during their entire shelf life. In addition, the Regulation sets down certain sanitation criteria regarding the production process.
Implications for Third Countries Exporting to the EU:

The European Commission developed a Guidance Document that addresses the key questions related to EU imports requirements. Food business operators will find the information they need regarding this new Hygiene regime on their businesses.

C) Feed Hygiene:

Contaminated feed has been responsible for many food crises. Council Regulation 183/2005 aims to ensure feed safety at all stages, including primary production. Effective January 1, 2006, it includes mandatory registration of feed growers, processors, packers and distributors with their competent authority. Note: in the U.S., it is the FDA and NOAA. However, in the absence of specific implementing rules concerning third countries, the existing rules on EU imports apply.

D) Food and Feed Controls:

The Food and Feed Regulation on Official Controls (Regulation 2017/625) establishes a harmonized EU controls system that includes: food and feed safety and animal health and welfare standards. Third countries must guarantee that products intended for the EU market meet the necessary standards. This section does not include animal welfare controls unless there are explicit animal welfare provisions in specific bilateral agreements.

Questions & Answers on Food Controls.
VI. Which Certificate for Which Product?

<table>
<thead>
<tr>
<th>PRODUCT</th>
<th>CERTIFICATE-YES OR NO?</th>
<th>RELEVANT LEGISLATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fishery and Aquaculture Products (Including wild raw off scallops)</td>
<td>YES</td>
<td>• Decision 2006/199/EC</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Regulation 1012/2012</td>
</tr>
<tr>
<td>Aquaculture Products - Live Fish</td>
<td></td>
<td>• Regulation 1251/2008</td>
</tr>
<tr>
<td>• Eggs, gametes - for farming</td>
<td>YES</td>
<td></td>
</tr>
<tr>
<td>• For consumption</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aquaculture Products - Mollusks</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Eggs, gametes for growth, fattening, or relaying</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td>• For consumption</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bivalve Mollusks</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

A) Fishery Products:

Since June 15, 2011, shipments of fishery and aquaculture products must be accompanied by a health certificate. U.S. exporters should consult the NOAA Seafood Inspection Program web site. This health certificate is covered by Commission Decision 2006/199/EC -for public health- and Regulation 1012/2012 for animal health. This health certificate is valid for both fishery and aquaculture products. Wild roe-off scallops are considered as fishery products and must be accompanied by a fishery product health certificate.

B) Aquaculture Products:

The relevant EU legislation regarding aquaculture products is Regulation 1251/2008 and its amendments.

C) Live Bivalve Mollusks:

Per Commission Decision 2009/951, imports of U.S. bivalve mollusks, in whatever form, are not permitted into the EU since July 2010.

Depending on current discussion between the U.S. Food & Drug Administration and the European Commission, transatlantic shellfish trade may resume by 2021. For more information contact CSEU-NOAA Office.
VII. Fishmeal – Fish oil

A) Fishmeal:

A certificate “for processed animal protein not intended for human consumption,” according to the model listed in Regulation 1069/2009 and its implementing Regulation 142/2011 (Chapter 1 certificate), must accompany U.S. exports of fishmeal.

For U.S. exporters, NOAA issues this certificate. Fishmeal exports intended for the EU must be produced by EU-approved establishments. The list of U.S. approved animal by-products establishments can be found through the link: https://webgate.ec.europa.eu/sanco/traces/output/US/ABP-PET_US_en.pdf

B) Fish Oil:

Effective April 30, 2009, the amended Hygiene Legislation requires that fish oil intended for human consumption must meet the requirements for “regular” fishery products. As such, U.S. fish oil shipments to the EU must be produced in EU-approved establishments and accompanied by the same public health certificate as the certificate for fishery products. Fish oil products containing plant or other ingredients (wheat…) in quantity are to be considered composite products. Regulation 28/2012 is the framework Regulation covering composite products. Under this Regulation, establishments wishing to send composite products do not need registration and must use the certificate described in Annex I of the Regulation. Due to legal complexities regarding this specific certificate, no U.S. Federal Agency (NOAA, FDA, USDA, FSIS, APHIS…) can issue such a document yet.

However, NOAA can issue a fishery product certificate for flavored fish oil and fish oil containing plant-based antioxidants for human consumption. These products are not considered to be composite and must come from EU approved establishments.

Shipments of fish oil not intended for human consumption are controlled by different legislations and must be accompanied by a certificate according to the regulation described in Regulation 142/2011 - Chapter 9 certificate - mentioned above.

For a complete overview of fish oil import requirements into the EU, see link below: https://ec.europa.eu/food/sites/food/files/animals/docs/bips_guidance_fish-oil_en.pdf
VIII. Duties and Trade Measures

A) Background:

All EU fish tariffs were consolidated under the Tokyo Round of GATT. The average EU duty for Chapters 3, 1604 and 1605 is 17.2%, one of the highest in the world. The tariff range is from 0% - live eels, to 25% - canned mackerel, bonito and anchovies. The primary legislation covering tariffs is Commission Implementing Regulation 2019/1776. However, the EU provides different mechanisms to reduce duties. It claims that its overall tariff average is then reduced to around 3 to 4%: An overall duty-free scheme applies to Africa-Caribbean-Pacific (ACP) countries, signatories of the Lomé Convention, for all seafood products imported into the EU from ACP countries. The Generalized System of Preferences (GSP), which applies to developing countries, applies to all seafood products in Chapter 3. Products are classified according to categories, e.g., sensitive, semi-sensitive and very sensitive.

B) Tariffs Quotas:

Tariffs Suspensions have been suspended since January 2013 and replaced by Autonomous Tariff Quotas (ATQs) that are granted annually for a period of for 3 years. These tariffs quotas are covered by Council Regulation 2018/1977 for the period ending December 31, 2020.

Autonomous quotas are opened on an annual basis. Each product or group of products is subject to a quantitative limit. The quota remains opened until the limit is reached. Quantities and reduced duties may change every year depending on Member States’ demands usually based on national industry requirements. Compromises are reached usually at Ministerial level.

For questions on a specific duty rate, consult the following web site: http://ec.europa.eu/taxation_customs/dds2/taric/taric_consultation.jsp?Lang=en&Screen=0&redirectionDate=20110203

IX. How Do I Label My Seafood Product?

A) Legislative Background:

Various crises within the food chain, such as Foot & Mouth disease, BSE, horsemeat misidentification incidents and detection of heavy metals in a product, have reinforced the critical need for information, communication and transparency for consumers from the producers, processors, and marketers.

The three primary labeling Regulations are:

- Regulation 1169/2011 on the provision of food information to consumers.

Food manufacturers must indicate the source allergen on the label, if it is used as an ingredient at any level in pre-packed foods. Directive 2006/142/EC adds “mollusks and products thereof” to the list of potential allergens.

All new EU Regulations are based to ensure the consumer’s confidence and safety so that “the consumer will not be misled by any product or packaging.”

For sanitary purposes and to allow traceability of seafood products, EU legislation requests that all outer and inner packages bear:

1. The country of origin,
2. The commercial denomination of the products, and
3. The approval number of the establishment of origin.

Regulation 853/2004 (Annex II, point 7) requires that products intended for the final consumer such as canned products, must include the FDA approval number of the U.S. packer/processor/manufacturers as well as their address or that of the EU seller.

However, U.S. exporters should pay specific attention to article 5 of Commission Decision 2006/199 regarding products in bulk and those intended for further processing that introduces derogation to this rule.

Finally, Regulation 853/2004 (Annex II, paragraph 11) allows for minimal labeling instead of normal labeling requirements: “For products of animal origin that are placed in transport containers or large packages and are intended for further processing, handling, wrapping or packaging in another establishment, the mark may be applied to the external surface of the container or packaging.”

Those two items must be written or printed “indelibly.” The most desirable way would be to have them pre-printed on packages/cartons. In instances where stick-on labels may be used, they must not be easily destructible or removable. Labels must be in a language “easily understandable” by users and at least in one of the official languages of the country of final destination (distribution). Labels may be in several languages.

Commission Regulation 2001/2065/EC identifies specific requirements for the labeling of fishery and aquaculture products intended for the retail sector. This Regulation only concerns products from Chapter 3 of the Tariff Harmonized System, and not products from...
Chapter 16, for example, canned products. This legislation has been amended by Regulation 1379/2013.

Since December 13, 2014, there are new requirements for products intended to retail and mass caterers.

The required information on the label of all fishery and aquaculture products for sale at retailers and mass caterers is:

<table>
<thead>
<tr>
<th>Commercial Name of Species</th>
<th>Production Method</th>
<th>Fishing Gear</th>
<th>Catch Area</th>
</tr>
</thead>
</table>

The commercial name of the species. The Latin name is not mandatory on the label except if required by your client. The EC has established a list of applicable commercial names. This list is visible on DG Mare’s website. (Note: the Latin name is required on other documentation such as the health certificate)

The production method, e.g., aquaculture or fishery product. The appropriate language is: “caught in...”; “caught in fresh water;” “farmed;” or “cultivated.” However, it is up to Member States whether this information is required when the commercial designation and the area of capture make it obvious that the fish was caught at sea.

The fishing gear: The new legislation gives the choice of seven fishing gears to be included on labels. In case of multiple fisheries, operators can put several fishing gears on the same label.

Operators that want to add additional voluntary information on the type of fishing gear can do so. Annex III of Regulation 1379/2013 provides a list of additional info for each type of gear.

Seines
Trawls
Gillnets and similar nets
Surrounding nets and lift nets
Hooks and lines
Dredges
Plots and traps
**The catch area.** Products caught at sea must identify the area of capture, which is taken from the FAO list. However, only the general area must be mentioned, e.g., Pacific Ocean. Products caught in fresh water require a reference to the Member State or third country of origin of these products. Farmed products must reference the Member State or third country in which the product underwent final stage of development. Operators may well choose to provide additional information regarding the area.

To ensure exact traceability at all stages of the marketing process, fisheries and aquaculture products must be accompanied by a document indicating the information described above as well as the Latin name of the products. The document concerned can be the invoice.

Since January 2014, all seafood sold at RETAIL outlets must have nutritional information on the package. Regulation 1169/2011 identifies the minimum information that must be included on labels of products intended for retail or mass caterers. Exporters should pay specific attention to Article 9, the following articles as well as all annexes of the Regulation.

Other Regulations regarding ingredients, allergens and guidelines for the implementation of labeling legislation can be downloaded at: http://ec.europa.eu/food/food/labellingnutrition/foodlabelling/index_en.htm.

It is important to note that some Member States as well as countries that are part of the European Economic Area (EEA) may have additional requirements for seafood labeling. For further information on labeling, contact our office at the U.S. Mission to the European Union.
B) Specific Labeling Examples:

**Fresh, Chilled Products:**
1. Species (commercial & scientific name)
2. Country of Origin- Roman Letters, min. 2 cm
3. Presentation- Whole, Gutted, Fillet, etc.
4. Best Before Date- Not Mandatory per EU Legislation but Requested by Most Member States
5. Freshness, Grade & Size Category (for Species with Common Standards, min. 5 cm).
6. Net Weight in Kilograms (Kg)
7. Date of Grading and Dispatch
8. Name & Address (city + state + "FDA approval #" of Processor/Packer)
9. Freshness Grading is Only for Whole/Gutted Fresh Fish

**Live Bivalve Mollusks:**
1. Species- Common Name and Latin Name
2. Country of Dispatch
3. Date of Wrapping - at least Day and Month
4. Date of Durability or "These Animals Must Be Alive When Sold"
5. Net Weight - Kg.
6. Identification of the Dispatch Center by Its Approval Number
7. Name and Address – City & State of Packer + "FDA Approval #"
8. Interstate Certified Shellfish Shipper #

**Canned Products:**
1. Name of Product
2. Country of Origin
3. Net Weight in Grams - or Liter for Liquid Products
4. Net Drained Weight- in Case of Solid Packed in a Usually- Not Consumed Liquid
5. List of Ingredients (added water is an ingredient)
6. Date of Minimum Durability - Year
7. Special Storage Conditions or Conditions of Use
8. Instructions for Use- If Not Obvious
9. Name and Address of Manufacturer, or EU Seller
10. "FDA Approval #" of Packer or Manufacturer/Processor
### Frozen Products:

1. Species (commercial & scientific name) followed by "Frozen"
2. Country of Origin
3. Presentation may be included with the name of the species
4. Net Weight in Kg
5. List of Ingredients - unless it is only fish
6. Date of Minimum Durability - month/year, or "Best Before" date
7. Freezing date
8. Special Storage Conditions - "to be maintained at -18°C"
9. Instructions for Use - if not obvious, include "Do Not Freeze Again Once Thawed"
10. Name and Address of Manufacturer or EU Seller
11. "FDA Approval #" of Packer (CFN or FEI) or Processor
12. Lot # - must begin with "L" or the word "LOT." Not always mandatory. The lot # is defined by the processor in order to trace a product history in case of problem. (It can be the production date).

### Deep-Frozen Products:

1. Same Requirements for Frozen Products, plus...
2. Freezing Date
3. Storage Conditions and Maximum Period of Storage:
   - Between 0 and 5°C: 1 day
   - "**", or between -5 and 0°C: 1 week
   - "***", or between -12 and -6°C: 1 month
   - "****", or at least at -18°C: up to the best before date

### Lot # Example:

L8110B15 may mean
- L = Lot
- 8 = 1998
- 110 = Day of Production
- B15 = Production Line Number
X. Other legislation

In addition to the above-mentioned legislation, the EU sets various requirements for a wide range of issues. This includes legislation on:

- **Additives**, colorings, **flavorings** and sweeteners allowed within the EU. Per Directive 95/2/EEC (amended several times), additives such as STP - E338 to E 450 - are not allowed in fresh scallops, only in frozen and deep-frozen scallops.

- **Traceability of foodstuffs**.

- **Packaging materials**: this legislation is important and relates to the characteristics of materials to not transfer substances to foodstuffs in quantities that may be harmful to human health or change organoleptic properties.

- **Sport caught fish**: [Commission Delegated Regulation 2019/2122](#) identifies weight limits under which there is no need for health certificate for the import of fish for personal consumption. This limit has been raised from 1 kg to 20 kg. Article 7, paragraph h.

XI. Illegal, Unreported, and Unregulated (IUU) Legislation

In 2008, the EU adopted [Council Regulation (EC) 1005/2008](#) aimed at eliminating Illegal, Unreported & Unregulated (IUU) fishing. Since January 1, 2010, all third countries wishing to export seafood to the EU must provide a catch certificate.

**This catch certificate is required in addition to all other sanitary documentation.**

This Regulation has been amended by [Regulation 86/2010](#) and [Regulation 202/2011](#). These Regulations amend the list of products excluded by the scope of the catch certification system and identify specific agreements between the EC and third countries, including the U.S.

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**Catch Certificates are required for most products of Chapter 3 and 16 of the Combined Nomenclature.**  
For example, fish oil bearing HS code 1504 is **not** subject to Catch Certification.
Implementing measures, list of products excluded by the IUU legislation, list of competent Member States (MS) authorities, and FAQs are all found on the DG Mare website below: https://ec.europa.eu/fisheries/cfp/illegal_fishing/

NOAA signed an agreement with the EC that allows for a U.S. specific catch certificate. NOAA is responsible for the issuance of both sanitary and catch certificates. U.S. exporters will find the necessary information regarding these catch certificates, as well as FAQs on the NOAA SIP web site. As of March 1, 2020, NOAA will implement a new catch certificate that will include catching vessels.

Diagram of documents required for direct and indirect seafood exports to the EU (courtesy Linda Chaves):
Annex I: CERTIFICATION CHECKLIST

Establishment Listing:

- Is your establishment on the EU list of U.S. approved establishments?
- If yes, does it correspond to the one mentioned on the health certificate?
- No typos? Inversion of numbers?
- Is your third country supplier, if any, an EU approved establishment?
- Is there any restriction on products from that specific third country?

Date of certificate:

- Is your certificate dated prior to the bill of lading? If not, DON’T SHIP!

Approval number:

- Don't forget to mention your approval number on all (outer & inner) packages, irrespective of the final use of your product.
- Does the approval number on the health certificate match the one mentioned on your label?

Country of origin:

- Should always be USA.

Identification of products:

- Use the complete description of your products according to WCO nomenclature: “raw dressed” is not an accurate description of a product.

Transport info:

- Container and seal numbers not mandatory but highly recommended.

Label:

- Is your label complete?
- Does it match with box I.28 of your health certificate?
Annex II

XII. Points of contact

N.O.A.A. – National Marine Fisheries Service

<table>
<thead>
<tr>
<th>Seafood Inspection Program:</th>
<th>Phone: (301) 427-8300</th>
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<tbody>
<tr>
<td>Steven Wilson</td>
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<tr>
<td>Robert Downs</td>
<td><a href="mailto:Robert.downs@noaa.gov">Robert.downs@noaa.gov</a></td>
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**Regional Offices:**

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<th>North-East: Inspection</th>
<th>Lawrence Biondo</th>
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<td>Phone: (978) 281-9228</td>
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<th>South-East Inspection</th>
<th>Brian Vaubel</th>
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<td>Fax: (727) 570-5387</td>
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<tr>
<th>South-West Inspection</th>
<th>Laurice Churchill</th>
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<td>Phone: (562) 388-7346</td>
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<td>Fax: (562) 388-7353</td>
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<tr>
<th>North-West Inspection</th>
<th>Eric Staiger</th>
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<tr>
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<td>Phone: (206) 526-4259</td>
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<td>Fax: (206) 526-4264</td>
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**Food and Drug Administration (FDA)**

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<th>Office of Seafood (Washington, DC):</th>
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<td>Fax: (202) 418-3196</td>
</tr>
<tr>
<td>Johnny Braddy</td>
<td><a href="mailto:Johnny.Braddy@fda.hhs.gov">Johnny.Braddy@fda.hhs.gov</a></td>
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**Food Guidance Document.**
Annex III - Useful links

EU List of U.S. FDA approved seafood producers/freezing & factory vessels/cold storage:
https://webgate.ec.europa.eu/sanco/traces/output/non_eu_listsPerCountry_en.htm#

FDA list of approved shellfish growers/shuckers/packers:
https://www.fda.gov/food/federalstate-food-programs/interstate-certified-shellfish-shippers-list

EU Official Journal:

DG SANTE - EU food safety legislation:
http://ec.europa.eu/food/food/index_en.htm

EU Tariffs database:
http://madb.europa.eu/madb/euTariffs.htm

DG Mare:
http://ec.europa.eu/fisheries/index_en.htm

European Food Safety Authority (EFSA)
http://www.efsa.europa.eu/

UK Seafish Organization
http://www.seafish.org
For More Information

The U.S. Commercial Service at the U.S. Mission to the European Union, can be contacted via email at: Stéphane Vrignaud, at stephane.vrignaud@trade.gov; Phone: +32(2) 811-5831; Fax: +32(2) 811 5151; or visit our website: http://www.export.gov/europeanunion.

The U.S. Commercial Service – Your Global Business Partner

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