supplement those otherwise maintained under the EAR for that particular country. This part does not address license requirements and licensing policies for controls implementing U.N. sanctions. CCL entries containing items subject to U.N. sanctions will refer the exporter to part 746 of the EAR, Embargoes and Other Special Controls, for any supplemental controls that may apply to exports and re-exports involving these countries.

(c) Exports and reexports involving Cuba and Iran. This part does not cover license requirements and licensing policies that apply to exports and reexports to embargoed destinations (Cuba and Iran). These comprehensive embargoes cover a broader range of items than those reflected in the CCL. If you are exporting or reexporting to any of these destinations, you should first review part 746 of the EAR, Embargoes and Other Special Controls.

(d) Anti-terrorism Controls on Cuba, Iran, North Korea, Sudan, and Syria. Commerce maintains anti-terrorism controls on Cuba, Iran, North Korea, Syria, and Sudan under section 6(a) of the Export Administration Act. Items controlled under section 6(a) to Iran, Syria, Sudan, and North Korea are described in §§ 742.8, 742.9, 742.19, and 742.19, respectively, and in Supplement No. 2 to part 742. Commerce also maintains controls under section 6(j) of the EAA to Cuba, Iran, North Korea, Sudan, and Syria. Items controlled to these countries under EAA section 6(j) are also described in Supplement 2 to part 742. The Secretaries of Commerce and State are required to notify appropriate Committees of the Congress 30 days before issuing a license for an item controlled under section 6(j) to Cuba, North Korea, Iran, Sudan, or Syria. If you are exporting or reexporting to Cuba, Iran, or North Korea, you should review part 746 of the EAR, Embargoes and Other Special Controls.

(e) End-user and end-use based controls. This part does not cover prohibitions and licensing requirements for exports of items not included on the CCL that are subject to end-use and end-user controls: certain nuclear end-uses; certain missile end-uses; certain chemical and biological weapons end-uses; certain naval nuclear propulsion end-uses; certain activities of U.S. persons; and certain exports to and for the use of certain foreign vessels and aircraft. Licensing requirements and policies for these exports are contained in part 744 of the EAR.

(f) Overlapping license policies. Many items on the CCL are subject to more than one type of control (e.g., national security (NS), missile technology (MT), nuclear nonproliferation (NP), regional stability (RS)). In addition, applications for all items on the CCL, other than those controlled for short supply reasons, may be reviewed for missile technology (see § 742.5(b)(3) of this part), nuclear nonproliferation (see § 742.3(b)(2) of this part), or chemical and biological weapons (see § 742.2(b)(3) of this part), if the end-use or end-user may be involved in certain proliferation activities. Finally, many multilaterally controlled items are reviewed for anti-terrorism reasons if they are destined for a terrorism-supporting country (see paragraph (d) of this section). Your application for a license will be reviewed under all applicable licensing policies. A license will be issued only if an application can be approved under all applicable licensing policies.


§ 742.2 Proliferation of chemical and biological weapons.

(a) License requirements. The following controls are maintained in support of the U.S. foreign policy of opposing the proliferation and illegal use of chemical and biological weapons. (See also § 742.18 of this part for license requirements pursuant to the Chemical Weapons Convention).

(1) If CB Column 1 of the Country Chart (Supplement No. 1 to part 738 of the EAR) is indicated in the appropriate ECCN, a license is required to all destinations, including Canada, for the following:

(i) Human pathogens, zoonoses, toxins, animal pathogens, genetically modified microorganisms and plant pathogens identified in ECCNs: 1C351, 1C352, 1C353, 1C354 and 1C360; and
(ii) Technology (ECCNs 1E001 and 1E351) for the production and/or disposal of microbiological commodities described in paragraph (a)(i) of this section.

(2) If CB Column 2 of the Country Chart (Supplement No. 1 to part 738 of the EAR) is indicated in the appropriate ECCN, a license is required to all destinations except countries in Country Group A:3 (see Supplement No. 1 to part 740 of the EAR) (Australia Group members) for the following:

(i) Chemicals identified in ECCN 1C350 (precursor and intermediate chemicals used in the production of chemical warfare agents).

(A) This license requirement includes chemical mixtures identified in ECCN 1C350.b, .c, or .d, except as specified in License Requirements Note 2 to that ECCN.

(B) This licensing requirement does not include chemical compounds created with any chemicals identified in ECCN 1C350, unless those compounds are also identified in ECCN 1C350.

(C) This licensing requirement does not apply to any of the following medical, analytical, diagnostic, and food testing kits that consist of pre-packaged materials of defined composition that are specifically developed, packaged, and marketed for diagnostic, analytical, or public health purposes:

(1) Test kits containing no more than 300 grams of any chemical controlled by ECCN 1C350.b or .c (CB-controlled chemicals also identified as Schedule 2 or 3 chemicals under the CWC) that are destined for export or reexport to CWC States Parties (destinations listed in Supplement No. 2 to part 745 of the EAR). Such test kits are controlled by ECCN 1C350 for CB and CW reasons, to States not Party to the CWC (destinations not listed in Supplement No. 2 to part 745 of the EAR), and for AT reasons.

(2) Test kits that contain no more than 300 grams of any chemical controlled by ECCN 1C350.d (CB-controlled chemicals not also identified as Schedule 1, 2, or 3 chemicals under the CWC). Such test kits are controlled by ECCN 1C995 for AT reasons.

(iii) Technology (ECCN 1E001) for the production and/or disposal of chemical precursors described in ECCN 1C350.

(iv) Technology (ECCNs 1E001 and 1E350) involving the following for facilities designed or intended to produce chemicals described in 1C350:

(A) Overall plant design;

(B) Design, specification, or procurement of equipment;

(C) Supervision of construction, installation, or operation of complete plant or components thereof;

(D) Training of personnel;

(E) Consultation on specific problems involving such facilities.

(v) Technology (ECCNs 1E001 and 1E351) for the production and/or disposal of chemical precursors described in ECCN 1C350.

(vi) Equipment and materials identified in ECCN 2B350 or 2B351 on the CCL, chemical detection systems controlled by 1A004.c for detecting chemical warfare agents and having the characteristics of toxic gas monitoring systems described in 2B351.a, and valves controlled by ECCN 2A226 or ECCN 2A292 having the characteristics of those described in 2B350.g, which can be used in the production of chemical weapons precursors or chemical warfare agents.

(vii) Equipment and materials identified in ECCN 2B352, which can be used in the production of biological agents.

(viii) Dedicated software identified in ECCN 2D351 for the “use” of toxic gas monitoring systems and their dedicated detecting components controlled by ECCN 2B351.

(ix) Technology identified in ECCN 2E001 for the “development” of software controlled by ECCN 2D351.

(x) Technology identified in ECCN 2E001, 2E002, or 2E301 for:

(A) The development, production, or use of items controlled by ECCN 2B350, 2B351, or 2B352; or

(B) The development or production of valves controlled by ECCN 2A226 or 2A292 having the characteristics of those described in ECCN 2B350.g.
(xi) Technology identified in ECCN 2E201 or 2E290 for the use of valves controlled by ECCN 2A226 or 2A292 having the characteristics of those described in 2B350.g.

(3) If CB Column 3 of the Country Chart (Supplement No. 1 to part 738 of the EAR) is indicated in the appropriate ECCN, a license is required to Country Group D:3 (see Supplement No. 1 to part 740 of the EAR) for medical products identified in ECCN 1C991.d.

(4) A license is required to States not Party to the CWC (destinations not listed in Supplement No. 2 to Part 745 of the EAR), for mixtures controlled by 1C395.a and test kits controlled by 1C395.b.

(b) Licensing policy. (1) License applications for the items described in paragraph (a) of this section will be considered on a case-by-case basis to determine whether the export or reexport would make a material contribution to the design, development, production, stockpiling or use of chemical or biological weapons. When an export or reexport is deemed to make such a material contribution, the license will be denied. When an export or reexport is intended to be used in a chemical weapons or biological weapons program, or for chemical or biological weapons terrorism purposes, it is deemed to make a material contribution. The factors listed in paragraph (b)(2) of this section are among those that will be considered to determine what action should be taken on license applications for these items.

(2) The following factors are among those that will be considered to determine what action should be taken on license applications for the items described in paragraph (a) of this section:

(i) The specific nature of the end-use, including the appropriateness of the stated end-use;

(ii) The significance of the export and reexport in terms of its potential contribution to the design, development, production, stockpiling, or use of chemical or biological weapons;

(iii) The nonproliferation credentials of the importing country, including the importing country’s chemical and biological capabilities and objectives;

(iv) The extent and effectiveness of the export control system in the importing country and in any intermediary country through which the items being exported or reexported will transit or be transshipped en route to the importing country;

(v) The risk that the items will be diverted for use in a chemical weapons or biological weapons program, or for chemical weapons or biological weapons terrorism purposes;

(vi) The reliability of the parties to the transaction, including whether:

(A) An export or reexport license application involving any such parties has previously been denied;

(B) Any such parties have been engaged in clandestine or illegal procurement activities;

(C) The end-user is capable of securely handling and storing the items to be exported or reexported;

(vii) Relevant information about proliferation and terrorism activities, including activities involving the design, development, production, stockpiling, or use of chemical or biological weapons by any parties to the transaction;

(viii) The types of assurances or guarantees against the design, development, production, stockpiling, or use of chemical or biological weapons that are given in a particular case, including any relevant assurances provided by the importing country or the end-user;

(ix) The applicability of other multilateral export control or nonproliferation agreements (e.g., the Chemical Weapons Convention and the Biological and Toxin Weapons Convention) to the transaction; and

(x) The existence of a pre-existing contract.

(3) BIS will review license applications in accordance with the licensing policy described in paragraph (b)(1) of this section for items not described in paragraph (a) of this section that:

(i) Require a license for reasons other than short supply; and

(ii) Could be destined for the design, development, production, stockpiling, or use of chemical or biological weapons, or for a facility engaged in such activities.

(4) License applications for items described in paragraph (a) of this section,
when destined for the People's Republic of China, will be reviewed in accordance with the licensing policies in both paragraph (b) of this section and §742.4(b)(7).

(c) Contract sanctity. Contract sanctity dates are set forth in Supplement No. 1 to part 742. Applicants who wish that a preexisting contract be considered in reviewing their license applications must submit documentation sufficient to establish the existence of such a contract.

(d) Australia Group. The Australia Group, a multilateral body that works to halt the spread of chemical and biological weapons, has developed common control lists of items specifically related to chemical and biological weapons. Australia Group members are listed in Country Group A:3 (see Supplement No. 1 to part 740 of the EAR). Controls on items listed in paragraph (a) of this section are consistent with lists agreed to in the Australia Group.

(e) License application requirements and instructions. (1) Supplement No. 1 to part 748 of the EAR provides general instructions for completing license applications. When preparing applications for items controlled for chemical and biological reasons, pay particular attention to the instructions contained in paragraphs (e) and (f) of the Supplement that apply to entering "Quantity" and "Units," respectively, on license applications. Paragraphs (e) and (f) require that, if an item is licensed in terms of "$ value" (refer to the "Unit" paragraph within the appropriate ECCN), the unit of quantity commonly used in the trade must also be shown on the license application. In such cases, Section 750.7 of the EAR provides that the quantity of commodities authorized is limited by the total dollar value as shown on the approved license and not by the quantity specified thereon. Although the EAR do not place a specific limitation on quantity in such cases, the total quantity that may be exported or reexported is limited, to a significant degree, by the fact that the EAR do not provide a shipping tolerance for items licensed by "dollar value" (see Section 750.11(b)(1) of the EAR) and require that the "unit price" indicated on the license application reflect the fair market value of the items listed on the application (see paragraph (g) of Supplement No. 1 to part 748 of the EAR).

(2) Unique application and submission requirements for chemicals, medicinals, and pharmaceuticals are described in paragraph (a) of Supplement No. 2 to part 748 of the EAR.

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EDITORIAL NOTE: For Federal Register citations affecting §742.2, see the List of CFR Sections Affected, which appears in the Finding Aids section of the printed volume and on GPO Access.

§742.3 Nuclear nonproliferation.

(a) License requirements. Section 309(c) of the Nuclear Non-Proliferation Act of 1978 requires BIS to identify items subject to the EAR that could be of significance for nuclear explosive purposes if used for activities other than those authorized at the time of export or reexport. ECCNs on the CCL that include the symbol "NP 1" or "NP 2" in the "Country Chart" column of the "License Requirements" section identify items that could be of significance for nuclear explosive purposes and are therefore subject to licensing requirements under this part and under section 309(c) of the Nuclear Non-Proliferation Act of 1978. These items are referred to as "The Nuclear Referral List" and are subject to the following licensing requirements:

(1) If NP Column 1 of the Country Chart (Supplement No. 1 to part 738 of the EAR) is indicated in the appropriate ECCN, a license is required to all destinations except Nuclear Suppliers Group (NSG) member countries (Country Group A:4) (see Supplement No. 1 to part 740 of the EAR).

(2) If NP Column 2 of the Country Chart (Supplement No. 1 to part 738 of the EAR) is indicated in the applicable ECCN, a license is required to Country Group D:2 (see Supplement No. 1 to part 740 of the EAR) except India.

(3) Other nuclear-related license requirements are described in §§744.2 and 744.5 of the EAR.

(b) Licensing policy. (1) To implement the controls in paragraph (a) of this section, the following factors are among those used to determine what action should be taken on individual applications: