

FACT SHEETS

Fact Sheet: President Donald J. Trump Bolsters National Security and Strengthens U.S. Supply Chains by Imposing Tariffs on Patented Pharmaceutical Products

The White House

April 2, 2026

BOLSTERING NATIONAL SECURITY: Today, President Donald J. Trump imposed tariffs on patented pharmaceuticals and their ingredients under Section 232 of the Trade Expansion Act of 1962 to bolster American national security and public health.

- President Trump imposed a 100% tariff on patented pharmaceutical products and ingredients.
- The tariffs will come into effect in 120 days for certain large companies, and 180 days for smaller companies.
- Trade Deal Countries:
 - If a pharmaceutical product is from the European Union, Japan, Korea, or Switzerland and Liechtenstein, a 15% tariff will apply. If a pharmaceutical product is from the United Kingdom, a lower tariff will apply, subject to the recently concluded UK pharmaceutical agreement. *Onshoring and pricing agreements:*
 - For companies that enter into Most Favored Nation (MFN) pricing agreements with the Department of Health and Human Services (HHS) and onshoring agreements with the Department of Commerce, a 0% tariff will apply through January 20, 2029. For companies that only enter into onshoring agreements with the Department of Commerce, a 20% tariff will apply. The Department of Commerce and HHS will provide pathways for companies to enter into onshoring and MFN pricing deals with the U.S. Government.
 - *Generic pharmaceuticals:* Generic pharmaceutical products, biosimilars, and associated ingredients are not subject to tariffs at this time. This will be reassessed in one year.
- *Specialty pharmaceutical products:* Orphan drugs, drugs for animal health, and certain other specialty pharmaceutical products will be exempt, if they are from trade deal countries or meet an urgent public health need.

- The Proclamation establishes strong monitoring and enforcement mechanisms, including external audits and tariff increases on future and past imports.

STRENGTHENING AMERICAN SUPPLY CHAINS: President Trump recognizes that America must manufacture pharmaceutical products in order to be safe, secure, and healthy.

- President Trump imposed these tariffs following an extensive investigation conducted by the Secretary of Commerce under Section 232 of the Trade Expansion Act of 1962, as amended, to determine the effects on national security of imports of pharmaceuticals, pharmaceutical ingredients, and related products.
- The investigation found that patented pharmaceuticals and associated pharmaceutical ingredients are being imported into the United States in such quantities and under such circumstances as to threaten to impair our national security.
- The impending Section 232 tariffs have already spurred approximately \$400 billion in new investment commitments from U.S. and foreign pharmaceutical companies, which will be spent in the United States during President Trump's current term in office.
- A self-sufficient domestic manufacturing and industrial base for pharmaceutical products is vital for the ability to support national defense requirements and public health.
- Despite being the world leader in research and development for most innovative pharmaceuticals, the U.S. is heavily reliant on imports, threatening to limit U.S. access to life-saving medications in the event of global supply chain disruption.

BUILDING ON PROMISES KEPT TO PUT AMERICA FIRST: This action builds on President Trump's commitment to put America first, protect our national security, and strengthen American manufacturing across all sectors.

- In May 2025, President Trump signed an Executive Order to remove regulatory barriers and facilitate the restoration of a robust domestic manufacturing base for prescription drugs, including key ingredients and materials necessary to manufacture prescription drugs.
- In August 2025, President Trump signed an Executive Order to ensure American pharmaceutical supply chain resilience by filling the strategic active pharmaceutical ingredients reserve.
- President Trump's Administration has launched Section 232 investigations in adjacent sectors such as personal protective equipment, medical consumables, and medical equipment and devices, as well as robotics. These investigations will help ensure that harmful imports in any strategic sector do not compromise national security.

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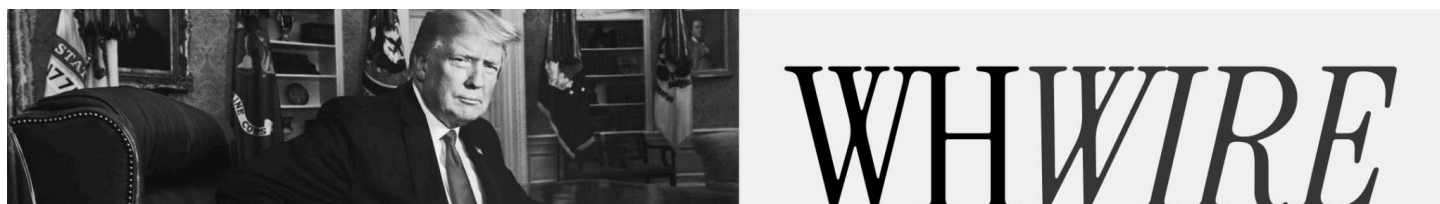
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PRESIDENTIAL ACTIONS

ADJUSTING IMPORTS OF PHARMACEUTICALS AND PHARMACEUTICAL INGREDIENTS INTO THE UNITED STATES

Proclamations

April 2, 2026

BY THE PRESIDENT OF THE UNITED STATES OF AMERICA

A PROCLAMATION

1. The Secretary of Commerce (Secretary) recently transmitted to me a report on his investigation into the effects of imports of pharmaceuticals and pharmaceutical ingredients on the national security of the United States under section 232 of the Trade Expansion Act of 1962, as amended, 19 U.S.C. 1862 (section 232). Based on the facts considered in that investigation, and taking into account the close relation of the economic welfare of the Nation to our national security and other relevant factors, see 19 U.S.C. 1862(d), the Secretary found and advised me of his opinion that pharmaceuticals and associated active pharmaceutical ingredients (APIs), including key starting materials, are being imported into the United States in such quantities and under such circumstances as to threaten to impair the national security of the United States.
2. The Secretary found that the present quantities and circumstances of imports of pharmaceuticals and pharmaceutical ingredients threaten to impair the national security and economy. Despite being the world leader in research and development (R&D) for most innovative pharmaceuticals (those that are typically patented and branded, as compared to generic pharmaceuticals or pharmaceuticals approved pursuant to section 505(j) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. 355(j)), the United States is heavily reliant on imports, threatening to limit United States access to life-saving medications in the event of global supply chain disruption due to geopolitical or economic disruption. According to the Food and Drug Administration, as of 2025, approximately 53 percent of patented pharmaceutical products distributed domestically are produced outside the country. The degree of import reliance is

significant at the API level with only 15 percent of patented APIs by volume domestically produced for the United States market.

3. The Secretary found that patented pharmaceuticals and associated pharmaceutical ingredients are essential to the United States' military and civilian healthcare. A self-sufficient domestic manufacturing and industrial base for pharmaceutical products is vital for the ability to support national defense requirements and maintain public health security during a national emergency or wartime. Patented pharmaceuticals are pivotal for treating cancer, rare diseases, autoimmune disorders, infectious diseases, and other critical health challenges. The Secretary further found that foreign government intervention has undermined the competitiveness of the United States patented pharmaceutical industry. This intervention has led to further dependence on foreign production of patented pharmaceuticals that have fragile supply chains.

4. In light of these findings, the Secretary recommended actions to adjust imports of patented pharmaceuticals and associated pharmaceutical ingredients, including continuing to negotiate onshoring agreements related to Most-Favored-Nation (MFN) pharmaceutical pricing agreements; imposing significant tariffs on pharmaceuticals and pharmaceutical ingredients, so that such imports will not threaten to impair the national security of the United States; and granting preferential treatment to those companies that commit to onshore production of pharmaceuticals and pharmaceutical ingredients.

5. After considering the Secretary's report, the factors in section 232(d) (19 U.S.C. 1862(d)), and other relevant factors and information, among other things, I concur with the Secretary's finding that pharmaceuticals and associated pharmaceutical ingredients are being imported into the United States in such quantities and under such circumstances as to threaten to impair the national security of the United States. In my judgment, and in light of the Secretary's report, the factors in section 232(d) (19 U.S.C. 1862(d)), and other relevant factors and information, I have also determined that it is necessary and appropriate to adopt a plan of action, as described below, to adjust such imports of pharmaceuticals and associated pharmaceutical ingredients so that such imports will not threaten to impair the national security of the United States.

6. I have decided to direct the Secretary and the Secretary of Health and Human Services to pursue negotiations of agreements or continue any current negotiations of agreements, such as agreements contemplated in section 232(c)(3)(A)(i) (19 U.S.C. 1862(c)(3)(A)(i)), to address the threatened impairment of the national security with respect to imported patented pharmaceuticals and associated pharmaceutical ingredients, with any party the Secretary and the Secretary of Health and Human Services deem appropriate, and to update me on the progress of such negotiations within 90 days of the date of this proclamation. Under current circumstances and in light of future requirements of the United States, this action is necessary and appropriate to address the threatened impairment of the national security.

7. I have determined that it is necessary and appropriate to impose a 100 percent *ad valorem* duty rate on the import of patented pharmaceuticals and associated pharmaceutical ingredients, as identified in Annex I to this proclamation, and except as otherwise provided in

this proclamation. Pharmaceutical products and ingredients that are subject to the section 232 zero tariff at this time are listed in Annex IV to this proclamation.

8. I have determined that it is necessary and appropriate that the *ad valorem* duty rate be 20 percent on imports of patented pharmaceuticals and associated pharmaceutical ingredients produced by companies that have plans, approved by the Secretary, to onshore production of such pharmaceuticals and pharmaceutical ingredients. The aforementioned 20 percent rate shall increase to 100 percent 4 years after the date of this proclamation.

9. I have further determined that it is necessary to implement pharmaceutical-related commitments in existing trade deals with the European Union, Japan, the Republic of Korea, and Switzerland and Liechtenstein jointly, as well as a future pharmaceutical-related deal with the United Kingdom (on which the United States and the United Kingdom have reached an agreement in principle as of December 1, 2025). These deals further United States economic and national security interests.

10. I further find that it is necessary and appropriate to impose no tariffs on imports of patented pharmaceuticals and associated pharmaceutical ingredients produced by companies that have fully executed agreements or are negotiating agreements with the Secretary and the Secretary of Health and Human Services regarding MFN pricing and onshoring of production and R&D of patented pharmaceuticals and pharmaceutical ingredients. Such agreements further United States economic and national security interests by making pharmaceuticals more accessible and affordable in the United States and by strengthening the domestic manufacturing base.

11. I have further determined not to adjust imports of generic pharmaceuticals and their associated ingredients, including biosimilar products, at this time. This determination includes purchases of generic pharmaceuticals and ingredients for the Strategic API Reserve. I find that such products should not be subject to section 232 tariffs at this time.

12. In my judgment, based on current circumstances as well as the future needs of the United States, the actions in this proclamation are necessary and appropriate to address the threatened impairment of the national security posed by imports of pharmaceuticals and pharmaceutical ingredients.

13. Section 232 authorizes the President to take action to adjust the imports of an article and its derivatives that are being imported into the United States in such quantities or under such circumstances as to threaten to impair the national security. Section 232 includes the authority to adopt and carry out a plan of action, with adjustments over time, to address the national security threat. This plan of action may include negotiations of agreements along with other actions to adjust imports to address the national security threat, including tariffs. If action under section 232 includes the negotiation of an agreement, such as one contemplated in section 232(c)(3)(A)(i) (19 U.S.C. 1862(c)(3)(A)(i)), the President may also take other actions he deems necessary to adjust imports to eliminate the threat that the imported article poses to the national security, including if such an agreement is not entered into within 180 days of the date of this proclamation, is not being carried out, or is ineffective. See 19 U.S.C. 1862(c)(3)(A).

14. Section 604 of the Trade Act of 1974, as amended (19 U.S.C. 2483) (section 604), authorizes the President to embody in the Harmonized Tariff Schedule of the United States (HTSUS) the substance of statutes affecting import treatment, and actions thereunder, including the removal, modification, continuance, or imposition of any rate of duty or other import restriction. NOW, THEREFORE, I, DONALD J. TRUMP, President of the United States of America, by the authority vested in me by the Constitution and the laws of the United States, including section 232, 19 U.S.C. 1862; section 604, 19 U.S.C. 2483; and section 301 of title 3, United States Code, do hereby proclaim as follows:

(1) The Secretary and the Secretary of Health and Human Services, and any senior official they deem appropriate, shall pursue or continue pursuing negotiations of agreements, as contemplated in section 232(c)(3)(A)(i) (19 U.S.C. 1862(c)(3)(A)(i)), to address the threatened impairment of the national security with respect to imported pharmaceuticals and pharmaceutical ingredients.

(2) I hereby ratify, and delegate to the Secretary the authority necessary to enter into, the company-specific tariff agreements listed in Annex II to this proclamation that the Secretary entered into prior to this proclamation. I also hereby delegate to the Secretary the authority to enter into and implement similar agreements in the future, as referenced in clause (1) of this proclamation. The Secretary is authorized to monitor and enforce these agreements as he deems appropriate, consistent with clause (6) of this proclamation and applicable law.

(3)(a) Imports of patented pharmaceuticals and associated pharmaceutical ingredients, as listed in Annex I to this proclamation, will be subject to a 100 percent *ad valorem* duty rate.

(b) The *ad valorem* duty rate for patented pharmaceuticals and associated pharmaceutical ingredients, as listed in Annex I to this proclamation, shall be 20 percent for products of companies that have, or that the Secretary assesses are likely soon to have (e.g., based on agreements in principle), onshoring plans approved by the Secretary. The aforementioned 20 percent rate shall increase to 100 percent on April 2, 2030.

(c) The *ad valorem* duty rate for patented pharmaceuticals and associated pharmaceutical ingredients, as listed in Annex I to this proclamation, shall be 15 percent for products of Japan, the European Union, the Republic of Korea, and Switzerland and Liechtenstein jointly, unless a lower rate applies under clause (3) of this proclamation. The tariff rate on patented pharmaceuticals and associated pharmaceutical ingredients for products of the United Kingdom shall be 10 percent and then reduce to zero to the extent required by any future agreement between the United States and the United Kingdom on pharmaceutical pricing. The Secretary shall publish a *Federal Register* notice should the rate for the United Kingdom be reduced to zero.

(d) The *ad valorem* tariff rate shall be zero for drugs and associated ingredients, where all approved indications are designated as orphan pursuant to the Orphan Drug Act, 21 U.S.C. 360aa *et seq.*, and its implementing regulations; nuclear medicines; plasma derived therapies; fertility treatments; cell and gene therapies; antibody drug conjugates; medical countermeasures related to chemical, biological, radiological, and nuclear threats; or other specialty

pharmaceutical products to be identified by the Secretary, as well as pharmaceutical products for animal health, provided that the Secretary, in consultation with the United States Trade Representative (Trade Representative) and the Secretary of Health and Human Services, determines that: (1) they are products of a jurisdiction that has a current or forthcoming trade and security framework agreement as referenced in Executive Order 14346 of September 5, 2025 (Modifying the Scope of Reciprocal Tariffs and Establishing Procedures for Implementing Trade and Security Agreements), or (2) they meet an urgent United States health need. The Secretary shall publish a *Federal Register* notice whenever he makes such a determination.

(e) For companies that are eligible for the tariff treatment outlined in clause (3)(b) of this proclamation, and that have entered into MFN pharmaceutical pricing agreements with the Secretary of Health and Human Services, the applicable *ad valorem* tariff rate for pharmaceuticals and associated pharmaceutical ingredients shall be zero until January 20, 2029. The Secretary shall apply this zero tariff rate to companies that he determines are likely to be eligible soon (e.g., because they have agreements in principle with the Secretary and the Secretary of Health and Human Services). For avoidance of doubt, this zero tariff rate shall also apply per the terms of the agreements listed in Annex II to this proclamation.

(f) The Secretary may increase the tariff rates referenced in clause (2) of this proclamation, and in clauses (3)(b) and (3)(e) of this proclamation, to address companies' failure to fulfill commitments under the relevant plans and agreements. The Secretary, in consultation with the Trade Representative, may increase the tariff rates referenced in clause (3)(c) of this proclamation to address foreign jurisdictions' failure to fulfill commitments under agreements with the United States. The Secretary shall publish a *Federal Register* notice when tariff rates are increased.

(4) The tariffs and tariff treatment imposed by this proclamation shall be effective with respect to goods entered for consumption, or withdrawn from warehouse for consumption, on or after 12:01 a.m. eastern daylight time on July 31, 2026, for the companies listed in Annex III to this proclamation and September 29, 2026, for other companies and shall continue in effect, unless such actions are expressly reduced, modified, or terminated.

(5) Generic pharmaceuticals and their associated ingredients shall not be subject to tariffs pursuant to section 232 at this time. Within 1 year of the date of this proclamation, the Secretary shall, in consultation with any senior executive branch officials the Secretary deems appropriate, inform the President of any circumstances that, in the Secretary's opinion, might indicate the need to take action to adjust the imports of generic pharmaceuticals and their associated ingredients.

(6) The Secretary, in consultation with the Secretary of Health and Human Services, shall establish criteria for onshoring plans referenced in clause (3)(b) of this proclamation, to be published in the *Federal Register*. All onshoring plans shall be subject to approval, monitoring, and enforcement by the Secretary. The Secretary shall require companies with qualifying onshoring plans to submit periodic reports to the Secretary regarding progress towards fulfilling onshoring milestones. The Secretary may require that such reports be audited by an external

auditing firm. In cases where the executive branch assesses that a company engaged in fraud or deliberately misled the United States Government with respect to onshoring commitments, the Secretary may reimpose tariffs discussed in this proclamation both prospectively and retroactively on imports from relevant companies, and he may impose other tariffs and penalties to the extent consistent with applicable law.

(7) If a product is subject to tariffs under this proclamation and Column 1 of the HTSUS (Column 1 Duty Rate), then the sum of the additional section 232 tariff imposed pursuant to this proclamation and the applicable Column 1 Duty Rate shall be equal to the applicable rate listed in clause (3) of this proclamation, unless the Column 1 Duty Rate is greater than the applicable rate listed in clause (3) of this proclamation, in which case only the Column 1 Duty Rate shall apply. This clause does not apply to the tariff treatment for products of the United Kingdom described in clause (3)(c) of this proclamation.

(8) If a product is subject to more than one rate of duty under this proclamation, then the lowest applicable rate shall apply.

(9) The Secretary, in consultation with the Chair of the United States International Trade Commission and the Commissioner of U.S. Customs and Border Protection (CBP), shall determine whether any modifications to the HTSUS or other administrative measures are necessary to effectuate or implement this proclamation or any actions taken pursuant to this proclamation. Any changes shall be published in a notice in the *Federal Register*.

(10) Drawback shall be available with respect to the duties imposed pursuant to this proclamation.

(11) Imports of United States-origin pharmaceutical products shall not be subject to the tariffs imposed by this proclamation at this time.

(12) To the extent permitted by applicable law, CBP may take any necessary or appropriate measure to administer the tariffs imposed or altered by this proclamation. Importers shall provide to CBP information necessary to carry out this proclamation.

(13) Any product described in clause (4) of this proclamation, except those eligible for admission as “domestic status” as described in 19 CFR 146.43, that is subject to a duty imposed by this proclamation and that is admitted into a United States foreign trade zone on or after the effective date of this proclamation, must be admitted as “privileged foreign status” as described in 19 CFR 146.41 and will be subject upon entry for consumption to any *ad valorem* rates of duty related to the classification under the applicable HTSUS subheading.

(14) The Secretary shall continue to monitor imports of patented and generic pharmaceuticals and pharmaceutical ingredients. The Secretary also shall, from time to time, in consultation with any senior executive branch officials the Secretary deems appropriate, review the status of such imports with respect to the national security. The Secretary shall inform me of any circumstances that, in the Secretary’s opinion, might indicate the need for further action by the President under section 232. The Secretary shall also inform me of any circumstance that, in the Secretary’s opinion, might indicate that the tariff imposed in this proclamation is no longer necessary.

(15) To the extent consistent with applicable law and the purpose of this proclamation, the Secretary, the Secretary of Health and Human Services, and the Secretary of Homeland Security are directed and authorized to take all actions that are appropriate to implement and effectuate this proclamation and any actions contemplated by this proclamation, including, consistent with applicable law, the issuance of regulations, rules, guidance, and procedures and the temporary suspension or amendment of regulations, within their respective jurisdictions, and to employ all powers granted to me under section 232.

(16) The Secretary, the Trade Representative, and the Secretary of Homeland Security may, consistent with applicable law, including section 301 of title 3, United States Code, redelegate any of these functions within their respective executive departments or agencies.

(17) Any provision of previous proclamations and Executive Orders that is inconsistent with this proclamation is superseded to the extent of such inconsistency. If any provision of this proclamation or the application of any provision of this proclamation to any individual or circumstance is held to be invalid, the remainder of this proclamation and the application of its provisions to any other individual or circumstance shall not be affected.

IN WITNESS WHEREOF, I have hereunto set my hand this second day of April, in the year of our Lord two thousand twenty-six, and of the Independence of the United States of America the two hundred and fiftieth.

DONALD J. TRUMP

ANNEXES I, II, III & IV

Related

Adjusting Imports of Timber, Lumber, and their Derivative Products into the United States

Presidential Actions, Proclamations | September 29, 2025

ADJUSTING IMPORTS OF SEMICONDUCTORS, SEMICONDUCTOR MANUFACTURING EQUIPMENT, AND THEIR DERIVATIVE PRODUCTS INTO THE UNITED STATES

Presidential Actions, Proclamations | January 14, 2026

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Annex I

**MODIFICATIONS TO
THE HARMONIZED TARIFF SCHEDULE OF THE UNITED STATES**

A. Effective with respect to goods entered for consumption, or withdrawn from warehouse for consumption, on or after 12:01 a.m. eastern time on July 31, 2026, subchapter III of chapter 99 of the Harmonized Tariff Schedule of the United States (“HTSUS”) is modified as follows:

1. The following new U.S. note 40 is inserted in numerical order:

“(a) Headings 9903.04.60–9903.04.69 provide the customs duty treatment of imported articles classifiable in one of the provisions of the HTSUS enumerated in subdivision (c) of this note.

These headings are mutually exclusive, such that an imported article will be subject to no more than one of these headings. Pharmaceutical articles, as defined in subdivisions (c)(i)–(iii) of this note, are subject to heading 9903.04.60 unless another of these headings applies.

(b) For imported articles subject to headings 9903.04.60–9903.04.68 that are eligible for special tariff treatment under any of the free trade agreements or preference programs listed in general note 3(c)(i) to the tariff schedule, the duties provided in these headings shall be collected in addition to any special rate of duty otherwise applicable under the appropriate tariff subheading. Goods for which entry is claimed under a provision of chapter 98 of the HTSUS and which are subject to the additional duties prescribed herein shall be eligible for and subject to the terms of such provision and applicable U.S. Customs and Border Protection (“CBP”) regulations. No claim for entry or for any duty exemption or reduction shall be allowed under a provision of chapter 99 of the HTSUS that may set forth a lower rate of duty or provide duty-free treatment, taking into account information supplied by CBP.

All antidumping, countervailing, or other duties and charges applicable to such goods shall continue to be imposed.

(c) The headings provided in subdivision (a) of this note and the defined terms of this subdivision apply to articles that are classifiable in the following provisions of the HTSUS:

2918.99.3000	2921.49.3800	2921.49.4300
2922.19.0900	2922.29.2700	2922.49.2600
2922.50.1400	2922.50.2500	2924.29.6250
2925.29.2000	2928.00.3000	2930.90.9235
2931.90.2200	2932.20.2000	2933.19.3500
2933.19.4500	2933.29.2000	2933.29.4500
2933.39.4100	2933.49.2600	2933.59.2100
2933.59.3600	2933.59.4600	2933.59.5300

2933.59.5900	2933.79.0800	2933.79.8500
2933.99.4600	2933.99.5300	2933.99.5590
2933.99.6100	2933.99.6500	2933.99.7000
2933.99.7500	2933.99.9000	2934.30.2300
2934.30.2700	2934.99.3000	2934.99.4700
2935.90.4800	2935.90.6000	2937.11.0000
2937.12.0000	2937.19.0000	2937.22.0000
2937.23.1010	2937.23.1050	2937.23.5010
2937.23.5020	2937.23.5050	2937.29.9040
2937.29.9050	2937.29.9095	2937.50.0000
2937.90.4500	2937.90.9000	2938.90.0000
2939.11.0000	2939.19.2000	2939.19.5000
2941.10.5000	2941.90.1050	2941.90.3000
2941.90.5000	2942.00.0500	3002.12.0040
3002.13.0010	3002.13.0090	3002.14.0010
3002.14.0090	3002.15.0011	3002.15.0091
3002.41.0000	3002.42.0000	3002.49.0050
3002.51.0000	3002.59.0000	3002.90.1000
3002.90.5220	3002.90.5250	3003.20.0000
3003.31.0000	3003.39.1000	3003.39.5000
3003.49.0000	3003.90.0120	3003.90.0140
3003.90.0180	3003.90.0190	3004.10.1010
3004.10.5010	3004.20.0010	3004.20.0083
3004.31.0000	3004.32.0060	3004.39.0010
3004.39.0055	3004.41.0000	3004.49.0005
3004.49.0010	3004.49.0020	3004.49.0030
3004.49.0040	3004.49.0050	3004.49.0060
3004.49.0070	3004.50.5005	3004.90.1000
3004.90.9201	3004.90.9206	3004.90.9208
3004.90.9210	3004.90.9211	3004.90.9215
3004.90.9216	3004.90.9225	3004.90.9236
3004.90.9243	3004.90.9246	3004.90.9249
3004.90.9251	3004.90.9252	3004.90.9253
3004.90.9260	3004.90.9263	3004.90.9267
3004.90.9268	3004.90.9270	3004.90.9271
3004.90.9273	3004.90.9276	

For the purposes of this note:

- (i) “Pharmaceutical articles” refers to imported articles classifiable in the provisions enumerated in this subdivision that are pharmaceutical products or that are ingredients (active pharmaceutical ingredients and key starting materials) classifiable in the provisions enumerated in this subdivision used to make pharmaceutical products.
- (ii) “Patented pharmaceutical articles” are pharmaceutical articles that are subject to a valid, unexpired U.S. patent and are listed in the U.S. Food and Drug

Administration's ("FDA") Approved Drug Products with Therapeutic Equivalence Evaluations ("Orange Book") or in the FDA's Lists of Licensed Biological Products ("Purple Book"); and ingredients (active pharmaceutical ingredients and key starting materials) for such articles.

- (iii) "Generic pharmaceutical articles" are FDA-approved pharmaceutical articles, and associated ingredients, that are not subject to a valid, unexpired U.S. patent and are off exclusivity. A generic pharmaceutical article is an active pharmaceutical ingredient or any component in a finished dosage form product that is used in a drug product or biosimilar biological product approved pursuant to a qualifying application; or a drug product or biosimilar biological product approved or licensed pursuant to a qualifying application. A qualifying application is: (a) an abbreviated new drug application submitted under section 505(j) of the Federal Food, Drug, and Cosmetic Act ("FDCA"); (b) a new drug application submitted under section 505(b)(2) of the FDCA that has been requested to be or that has been deemed therapeutically equivalent to a listed drug; (c) a biosimilar biologics application submitted under section 351(k) of the Public Health Services Act; or (d) an application for an authorized generic drug or authorized biological product, as those terms are described in section 505(t) of the FDCA and 42 U.S.C. § 1320f-1(e)(2)(B)(ii), provided that the products are imported by a generic or biosimilar manufacturer.

(d) Heading 9903.04.60 applies to patented pharmaceutical articles provided for in subdivision (c) of this note. For articles for which the applicable column 1 tariff rate is less than the additional duties provided by heading 9903.04.60, the sum of the column 1 duty rate and the additional *ad valorem* rate of duty shall be the rate of duty provided by heading 9903.04.60. For articles for which the applicable column 1 tariff rate is greater than the additional duties provided by heading 9903.04.60, then no additional duty is due pursuant to heading 9903.04.60.

(e) Heading 9903.04.61 applies to patented pharmaceutical articles imported for companies identified by the Secretary and imported before 12:01 am eastern time on September 29, 2026. The Secretary shall notify CBP of all such companies.

(f) Heading 9903.04.62 applies to patented pharmaceutical articles that are the product of Japan, of a member country of the European Union, of South Korea, of Switzerland, or of Liechtenstein that would otherwise be subject to the additional duties imposed under heading 9903.04.60. For articles for which the applicable column 1 tariff rate is less than the additional duties provided by heading 9903.04.62, the sum of the column 1 duty rate and the additional *ad valorem* rate of duty shall be the rate of duty provided by heading 9903.04.62. For articles for which the applicable column 1 tariff rate is greater than the additional duties provided by heading 9903.04.62, no additional duty is due pursuant to heading 9903.04.62.

(g) Heading 9903.04.63 applies to patented pharmaceutical articles that are the product of the United Kingdom that would otherwise be subject to the additional duties imposed under heading 9903.04.60.

(h) Headings 9903.04.64–9903.04.66 apply to patented pharmaceutical articles described in this subdivision. Any importer entering pharmaceutical articles under any of these headings shall provide any information that may be required, and in such form, as is deemed necessary by CBP to permit the administration of these headings.

(i) Heading 9903.04.64 applies to patented pharmaceutical articles imported for companies subject to an onshoring plan approved by the Secretary of Commerce in accordance with a process to be established in a Federal Register notice. The Secretary shall notify CBP of all such agreements.

(ii) Heading 9903.04.65 applies to patented pharmaceutical articles that meet the requirements of subdivision (h)(i) of this note and are imported for companies that have entered into a Most-Favored-Nation pharmaceutical pricing agreement with the Secretary of Health and Human Services. The Secretary of Commerce shall notify CBP of all such agreements.

(iii) Heading 9903.04.66 applies to drugs and associated ingredients for all approved indications that are designated as orphan pursuant to the Orphan Drug Act, 21 U.S.C. 360aa *et seq.* and its implementing regulations; nuclear medicines; plasma-derived therapies; fertility treatments; cell and gene therapies; antibody drug conjugates; medical countermeasures related to chemical, biological, radiological and nuclear threats; or other specialty pharmaceutical products identified by the Secretary of Commerce or pharmaceutical products for animal health imported from a jurisdiction that has a current or forthcoming trade and security framework or that meet an urgent U.S. health need. The Secretary shall publish a Federal Register notice when the conditions above are met and shall notify CBP of all such products.

(i) Heading 9903.04.69 applies to entries of articles that are classifiable under provisions of the HTSUS enumerated in subdivision (c) of this note but that are not pharmaceutical articles described in subdivisions (c)(i)–(iii) of this note.

2. The following new headings are inserted in numerical sequence, with the material in each new heading inserted in the columns of the HTSUS labeled “Heading/Subheading”, “Article Description”, “Rates of Duty 1-General”, “Rates of Duty 1-Special” and “Rates of Duty 2”, respectively:

Heading/ Subheading	Article Description	Rates of Duty		
		1		2
		General	Special	
“9903.04.60	Except as provided in heading 9903.04.61, patented pharmaceutical articles as provided for in subdivisions (c) and (d) of U.S. note 40 to this subchapter.....	100%	100%	The duty provided in the applicable subheading
9903.04.61	Patented pharmaceutical articles entered before 12:01 a.m. eastern time on September 29, 2026 as provided for in subdivisions (c) and (e) of U.S. note 40 to this subchapter.....	The duty provided in the applicable subheading	The duty provided in the applicable subheading	The duty provided in the applicable subheading
9903.04.62	Patented pharmaceutical articles that are the product of Japan, of a European Union member country, of South Korea, of Switzerland, or of Liechtenstein as provided for in subdivisions (c) and (f) of U.S. note 40 to this subchapter.....	15%	15%	The duty provided in the applicable subheading
9903.04.63	Patented pharmaceutical articles that are the product of the United Kingdom as defined in subdivisions (c) and (g) of U.S. note 40 to this subchapter.....	The duty provided in the applicable subheading + 10%	The duty provided in the applicable subheading + 10%	The duty provided in the applicable subheading
9903.04.64	Patented pharmaceutical articles subject to a qualifying onshoring plan, as provided for in subdivisions (c) and (h)(i) of U.S. note 40 to this subchapter.....	The duty provided in the applicable subheading + 20%	The duty provided in the applicable subheading + 20%	The duty provided in the applicable subheading
9903.04.65	Pharmaceutical articles subject to a qualifying onshoring plan and a Most-Favored-Nation pharmaceutical pricing agreement, as provided for in subdivisions (c) and (h)(ii) of U.S. note 40 to this subchapter.....	The duty provided in the applicable subheading + 0%	The duty provided in the applicable subheading	The duty provided in the applicable subheading

9903.04.66	Drugs and pharmaceutical articles for the specific uses provided in subdivisions (c) and (h)(iii) of U.S. note 40 to this subchapter.....	The duty provided in the applicable subheading + 0%	The duty provided in the applicable subheading	The duty provided in the applicable subheading
9903.04.67	Generic pharmaceutical articles, as provided for in subdivision (c) of U.S. note 40 to this subchapter.....	The duty provided in the applicable subheading	The duty provided in the applicable subheading	The duty provided in the applicable subheading
9903.04.68	Pharmaceutical products with an active pharmaceutical ingredient packaged in dosage form that is a product of the United States.....	The duty provided in the applicable subheading	The duty provided in the applicable subheading	The duty provided in the applicable subheading
9903.04.69	Articles as provided for in subdivision (i) of U.S. note 40 to this subchapter.....	The duty provided in the applicable subheading	The duty provided in the applicable subheading	The duty provided in the applicable subheading”

3. U.S. note 2(aa)(v) is modified by:

- a. redesignating subdivisions (a)–(k) as (2)–(12); and
- b. inserting “(1) patented pharmaceutical articles provided for in headings 9903.04.60, 9903.04.61, 9903.04.62, 9903.04.63, 9903.04.64, 9903.04.65 and 9903.04. 66;” in numerical order.

4. Heading 9903.03.06 is modified by deleting “, or medium- and heavy-duty vehicles or medium- and heavy-duty vehicle parts,” and inserting “, medium- and heavy-duty vehicles or medium- and heavy-duty vehicle parts, or patented pharmaceutical articles,” in lieu thereof.

B. Effective with respect to goods entered for consumption, or withdrawn from warehouse for consumption, on or after 12:01 a.m. eastern time on January 20, 2029, subchapter III of chapter 99 of the HTSUS is modified as follows:

- a. subdivision (h)(ii) of U.S. note 40 to this subchapter is deleted;
- b. subdivision (h)(iii) of U.S. note 40 to this subchapter is renumbered as subdivision (h)(ii);
- c. heading 9903.04.65 is terminated and deleted; and

- d. the article description of heading 9903.04.66 is modified by deleting “(h)(iii)” and inserting “(h)(ii)” in lieu thereof.
- C. Effective with respect to goods entered for consumption, or withdrawn from warehouse for consumption, on or after 12:01 a.m. eastern time on April 2, 2030, heading 9903.04.64 is amended by deleting “20%” each place that it appears and inserting “100%” in lieu thereof.

Annex II

Below are the company-specific agreements related to Section 232 tariffs on pharmaceutical products and ingredients that the Secretary of Commerce entered into prior to this proclamation.

1. AbbVie Inc. dated March 20, 2026
2. Amgen Inc. dated December 19, 2025
3. AstraZeneca Pharmaceuticals, LP dated March 20, 2026
4. Bristol Myers Squibb dated December 19, 2025
5. Boehringer Ingelheim Pharmaceuticals, Inc. dated December 19, 2025
6. Eli Lilly and Company dated February 23, 2026
7. EMD Serono, Inc. dated February 12, 2026
8. Genentech, Inc. dated December 19, 2025
9. Gilead Sciences, Inc. dated December 19, 2025
10. Merck Sharp & Dohme LLC dated February 11, 2026
11. Novartis Pharmaceuticals Corporation dated February 12, 2026
12. Novo Nordisk Inc. dated February 23, 2026
13. Sanofi S.A. dated December 19, 2025

Annex III

Annex III of this proclamation lists the companies whose tariff treatment shall be effective 120 days from the date of this proclamation.

1. AbbVie Inc.
2. Amgen Inc.
3. AstraZeneca Pharmaceuticals, LP
4. Bristol Myers Squibb
5. Boehringer Ingelheim Pharmaceuticals, Inc.
6. Eli Lilly and Company
7. EMD Serono, Inc.
8. Genentech, Inc.
9. Gilead Sciences, Inc.
10. GlaxoSmithKline LLC and ViiV Healthcare Company
11. Johnson & Johnson
12. Merck Sharp & Dohme LLC
13. Novartis Pharmaceuticals Corporation
14. Novo Nordisk Inc.
15. Pfizer Inc.
16. Regeneron Pharmaceuticals, Inc.
17. Sanofi S.A.

Annex IV

This annex includes the list of HTSUS codes that are not covered under the Annex I actions of this proclamation and are subject to the Section 232 action related to pharmaceuticals and pharmaceutical ingredients with a tariff rate of zero. Pursuant to section (4) of Proclamation 11012 of February 20, 2026, “Imposing a Temporary Import Surcharge to Address Fundamental International Payments Problems,” these codes are not subject to the surcharge imposed by Proclamation 11012.

2903.45.1000	2919.90.5010	2926.90.4801	2933.49.6000	2936.26.0000	3004.20.0076
2903.51.1000	2919.90.5050	2926.90.5010	2933.49.7000	2936.27.0000	3004.20.0080
2903.59.9000	2920.90.5100	2926.90.5050	2933.52.1000	2936.28.0000	3004.32.0010
2903.69.9000	2921.19.1100	2927.00.4000	2933.52.9000	2936.29.1000	3004.32.0040
2903.78.0000	2921.29.0010	2927.00.5000	2933.53.0000	2936.29.1610	3004.39.0020
2903.79.9030	2921.29.0020	2928.00.2500	2933.54.0000	2936.29.1620	3004.39.0030
2903.79.9070	2921.29.0030	2929.90.2000	2933.59.1000	2936.29.1630	3004.39.0040
2903.89.1500	2921.29.0055	2929.90.5015	2933.59.1500	2936.29.2000	3004.39.0045
2903.89.2000	2921.30.1000	2929.90.5018	2933.59.1800	2936.29.5020	3004.42.0000
2903.89.7010	2921.30.5000	2929.90.5020	2933.59.2200	2936.29.5030	3004.49.00
2903.89.7090	2921.42.9000	2929.90.5030	2933.59.7000	2936.29.5050	3004.50.1000
2903.92.0000	2921.46.0000	2929.90.5040	2933.59.8000	2936.90.0110	3004.50.2000
2904.99.4000	2921.49.4500	2929.90.5095	2933.59.8500	2936.90.0150	3004.50.3000
2905.29.9000	2921.49.5000	2930.20.2010	2933.59.9500	2937.21.0010	3004.50.4000
2905.39.9000	2921.59.8010	2930.20.2050	2933.69.6010	2937.21.0020	3004.50.5010
2905.59.1000	2921.59.8090	2930.20.9010	2933.69.6021	2937.21.0030	3004.50.5020
2905.59.9000	2922.11.0000	2930.20.9020	2933.69.6030	2937.21.0040	3004.50.5030
2906.19.5000	2922.14.0000	2930.20.9050	2933.69.6050	2937.23.1020	3004.50.5040
2906.29.6000	2922.19.2000	2930.30.6000	2933.72.0000	2937.23.2500	3004.60.0000
2907.29.9000	2922.19.3300	2930.90.2900	2933.79.1500	2937.23.50	3004.90.9203
2908.19.6000	2922.19.6000	2930.90.4910	2933.91.0010	2937.29.1000	3004.90.9204
2909.19.1800	2922.19.7000	2930.90.4920	2933.91.0050	2937.29.9020	3004.90.9205
2909.20.0000	2922.19.9000	2930.90.4950	2933.99.0100	2937.29.9030	3004.90.9207
2909.30.6000	2922.19.9610	2930.90.9208	2933.99.0200	2937.90.0500	3004.90.9209
2909.49.1000	2922.19.9619	2930.90.9210	2933.99.0500	2937.90.1000	3004.90.9212
2909.49.1500	2922.19.9690	2930.90.9212	2933.99.0600	2937.90.2000	3004.90.9213
2909.49.2000	2922.29.6100	2930.90.9222	2933.99.0800	2937.90.4000	3004.90.9217
2909.49.6000	2922.29.8110	2930.90.9225	2933.99.1100	2938.10.0000	3004.90.9218
2909.50.4010	2922.29.8190	2930.90.9231	2933.99.1200	2939.19.1000	3004.90.9219
2909.50.4050	2922.31.0000	2930.90.9251	2933.99.1600	2939.20.0010	3004.90.9220
2909.50.4500	2922.39.2500	2931.49.0005	2933.99.1701	2939.20.0050	3004.90.9222
2909.50.5000	2922.39.4500	2931.49.0008	2933.99.2200	2939.30.0000	3004.90.9223
2912.19.5000	2922.39.5000	2931.49.0010	2933.99.2400	2939.41.0000	3004.90.9224
2912.49.2600	2922.41.0010	2931.49.0015	2933.99.2600	2939.42.0000	3004.90.9226

2914.19.0000	2922.41.0090	2931.49.0020	2933.99.4200	2939.44.0000	3004.90.9227
2914.40.9000	2922.42.5000	2931.49.0025	2933.99.5100	2939.45.0000	3004.90.9228
2914.50.3000	2922.44.0000	2931.49.0055	2933.99.5510	2939.49.0300	3004.90.9229
2914.50.5000	2922.49.1000	2931.49.0080	2933.99.5520	2939.59.0000	3004.90.9230
2914.62.0000	2922.49.3000	2931.53.0000	2933.99.5530	2939.62.0000	3004.90.9232
2914.69.2100	2922.49.3700	2931.90.9010	2933.99.5800	2939.63.0000	3004.90.9233
2914.69.9000	2922.49.4910	2931.90.9021	2933.99.7900	2939.69.0000	3004.90.9234
2914.79.4000	2922.49.4915	2931.90.9025	2933.99.8210	2939.72.0000	3004.90.9235
2915.29.3000	2922.49.4950	2931.90.9029	2933.99.8220	2939.79.0000	3004.90.9237
2915.39.3100	2922.49.8000	2931.90.9030	2933.99.8290	2939.80.0010	3004.90.9238
2915.39.3500	2922.50.0700	2931.90.9035	2933.99.8500	2939.80.0050	3004.90.9239
2915.39.4700	2922.50.1000	2931.90.9040	2933.99.8900	2940.00.6000	3004.90.9240
2915.39.9000	2922.50.1100	2931.90.9052	2933.99.9701	2941.10.1000	3004.90.9241
2915.90.1010	2922.50.1300	2932.14.0000	2934.10.1000	2941.10.2000	3004.90.9242
2915.90.1050	2922.50.1700	2932.19.5100	2934.10.2000	2941.10.3000	3004.90.9244
2915.90.1400	2922.50.3500	2932.20.3000	2934.10.9000	2941.20.1000	3004.90.9245
2915.90.1810	2922.50.4000	2932.20.5010	2934.20.4000	2941.20.5000	3004.90.9247
2915.90.1890	2922.50.5000	2932.20.5020	2934.20.8000	2941.30.0000	3004.90.9248
2915.90.2000	2923.10.0000	2932.20.5030	2934.30.4300	2941.40.0000	3004.90.9250
2915.90.5010	2923.20.2010	2932.20.5050	2934.30.5000	2941.50.0000	3004.90.9254
2915.90.5050	2923.20.2050	2932.99.6100	2934.91.0000	2941.90.1010	3004.90.9255
2916.19.3000	2924.11.0000	2932.99.7000	2934.92.0000	2942.00.3500	3004.90.9256
2916.19.5000	2924.19.1110	2932.99.9010	2934.99.0100	2942.00.5000	3004.90.9257
2916.20.5000	2924.19.1120	2932.99.9090	2934.99.0300	3001.20.0000	3004.90.9258
2916.31.5000	2924.19.1130	2933.11.0000	2934.99.0500	3001.90.0110	3004.90.9261
2916.39.4600	2924.19.1150	2933.19.9000	2934.99.0600	3001.90.0150	3004.90.9262
2916.39.7900	2924.19.8000	2933.21.0000	2934.99.0700	3001.90.0165	3004.90.9264
2917.13.0030	2924.21.1600	2933.29.0500	2934.99.0800	3001.90.0195	3004.90.9265
2917.13.0090	2924.21.5000	2933.29.3500	2934.99.0900	3002.12.0010	3004.90.9266
2917.19.1000	2924.29.0100	2933.29.4300	2934.99.1100	3002.12.0020	3004.90.9269
2917.19.7020	2924.29.0300	2933.29.6000	2934.99.1200	3002.12.0030	3004.90.9272
2917.19.7050	2924.29.1000	2933.29.9000	2934.99.1500	3002.12.0090	3004.90.9274
2917.34.0110	2924.29.2300	2933.33.0100	2934.99.1600	3002.13.00	3004.90.9275
2917.34.0150	2924.29.2600	2933.34.0000	2934.99.1800	3002.14.00	3006.30.1000
2917.39.3000	2924.29.2800	2933.35.0000	2934.99.2000	3002.15.00	3006.30.5000
2918.11.5100	2924.29.3300	2933.37.0000	2934.99.3900	3002.49.0010	3006.60.0000
2918.13.5000	2924.29.5700	2933.39.0800	2934.99.4400	3002.90.5210	3006.70.0000
2918.16.5010	2924.29.6210	2933.39.1000	2934.99.7000	3003.10.0000	3006.93.1000
2918.16.5020	2924.29.6220	2933.39.2000	2934.99.9001	3003.41.0000	3006.93.2000
2918.16.5090	2924.29.7100	2933.39.2100	2935.50.0000	3003.42.0000	3006.93.5000
2918.19.6000	2924.29.7710	2933.39.2300	2935.90.0600	3003.90.0160	3006.93.6000

2918.19.9000	2924.29.7720	2933.39.2500	2935.90.1000	3004.10.1020	3006.93.8000
2918.22.1000	2924.29.7730	2933.39.2700	2935.90.1300	3004.10.1045	
2918.22.5000	2924.29.7790	2933.39.3100	2935.90.1500	3004.10.5048	
2918.23.3000	2924.29.8000	2933.39.6110	2935.90.2000	3004.10.5049	
2918.23.5000	2924.29.9500	2933.39.6120	2935.90.3000	3004.10.5055	
2918.29.2000	2925.12.0000	2933.39.6130	2935.90.3200	3004.10.5065	
2918.29.6500	2925.19.4200	2933.39.6191	2935.90.3300	3004.10.5075	
2918.29.7500	2925.19.9100	2933.39.9200	2935.90.4200	3004.20.0020	
2918.30.2500	2925.21.0000	2933.41.0000	2935.90.7500	3004.20.0030	
2918.30.3000	2925.29.6000	2933.49.0800	2935.90.9500	3004.20.0045	
2918.30.9000	2925.29.9000	2933.49.1000	2936.21.0000	3004.20.0055	
2918.99.4300	2926.30.1000	2933.49.1500	2936.22.0000	3004.20.0065	
2918.99.4700	2926.40.0000	2933.49.1700	2936.23.0000	3004.20.0070	
2918.99.5000	2926.90.1400	2933.49.2000	2936.24.0100	3004.20.0071	
2919.90.3000	2926.90.4300	2933.49.3000	2936.25.0000	3004.20.0072	