



99005105012001, 99005105012001

Apply for a WHO certificate for the export of medicinal products for human use without a marketing authorization in the exporting country

Heruntergeladen am 28.06.2025 https://fimportal.de/xzufi-services/307673659/L100012

Modul	Sachverhalt
Leistungsschlüssel	99005105012001, 99005105012001
Leistungsbezeichnung I	Apply for a WHO certificate for the export of medicinal products for human use without a marketing authorization in the exporting country
Leistungsbezeichnung II	Apply for a WHO certificate for the export of medicinal products for human use without a marketing authorization in the exporting country
Typisierung	3 - Bundesaufsichtsverwaltung: Regelung
Quellredaktion	Schleswig-Holstein
Freigabestatus Katalog	unbestimmter Freigabestatus
Freigabestatus Bibliothek	fachlich freigegeben (gold)
Begriffe im Kontext	





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Leistungstyp	Leistungsobjekt mit Verrichtung
Leistungsgruppierung	Arzneimittel (005)
Verrichtungskennung	Ausstellung (012)
SDG-Informationsbereich	
Lagen Portalverbund	Import und Export (2070200)
Einheitlicher Ansprechpartner	Nein
Fachlich freigegeben am	11.12.2024
Fachlich freigegen durch	Federal Ministry of Health (BMG)
Handlungsgrundlage	https://www.gesetze-im-internet.de/amg_1976/73a.h tml
Teaser	Are you based in Germany and would like to export a medicinal product that is not authorized in Germany for use in humans to a country outside the EU? Then you need a WHO certificate.
Volltext	To export medicinal products from Germany, you must apply for a WHO Certificate for Pharmaceutical Products (CPP). You need the WHO certificate in the importing third country for all regulatory situations relating to the local approval and import of your medicinal product. This may be necessary • in the context of marketing authorization applications • in the context of applications for renewal, extension, variation or review of a marketing authorization • for the import of medicinal products authorized in
	the exporting country Germany participates in the "World Health Organization (WHO) Certificate System on the Quality of Pharmaceutical Products in International Trade". Certificates under this system certify the marketability of the medicinal product in the country of origin and serve to facilitate the movement of medicinal products. The certificates are issued by the competent authority





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	of the federal state in which the medicinal product is manufactured (exporting country).
	Who submits the application?
	You can apply for the WHO Certificate for Pharmaceutical Products (CPP) as a
	manufacturing company
	company exporting the medicinal product
	If the
	• competent authority of the country of destination
	wishes to apply for the certificate, it requires written authorization from you.
	Additional services
	As part of the application, you can request additional services for the certificate if necessary. These can be, for example
	 Over-authentication by the Federal Office of Justice Legalization by the diplomatic or consular mission of the importing country in Germany Sealing with thread
	You can find out which additional services you require from the competent authority to which you wish to submit the certificate.
Erforderliche Unterlagen	
Voraussetzungen	
Kosten	Verwaltungsgebühr: 20€ Verwaltungsgebühr: 75€
Verfahrensablauf	You must apply for the WHO Certificate for Pharmaceutical Products (CPP) in writing using the application form. The form is written in German and in one other language. These are English, French or Spanish.





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	You must submit all the necessary documents with the application. If you do not have a marketing authorization for the medicinal product in Germany, you must prove that the competent authority of the country of destination has approved the import and that it is aware of the reasons for the lack of marketing authorization.
	When applying in writing, you must apply for a separate WHO certificate for each medicinal product and for each importing country.
	You must also observe the requirements of the competent national authority.
Bearbeitungsdauer	6 - 8 Woche(n)
Frist	There is no legal deadline.
weiterführende Informationen	https://www.auswaertiges-amt.de/de/service/fragenkat alog-node/12-apostille-ausl/606196
Hinweise	The following information is available:
	 The declaration of authorization status for a pharmaceutical product is not part of the application procedure. Batch certificates for pharmaceutical products are not part of the application procedure. Such a certificate is only applied for if state batch tests are prescribed for the product.
Rechtsbehelf	ObjectionAction before the administrative court within one month of notification
Kurztext	 WHO certificate (CPP) for the export of medicinal products for human use Issued without a marketing authorization in the exporting country WHO certificate (CPP) for the export of medicinal products: for human use without authorization in the exporting country Application for a certificate in accordance with the certificate system of the World Health Organization (WHO)





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	 WHO certificate (CPP) may be required for the export of a medicinal product or for regulatory purposes in a third country: in the context of marketing authorization applications in the context of applications for renewal, extension, variation or review of a marketing authorization for the import of medicinal products authorized in the exporting country Application procedure in writing using the appropriate form and online in some federal states Responsible: State authority in which the manufacturing or exporting company is based
Ansprechpunkt	
Zuständige Stelle	
Formulare	
Ursprungsportal	WHO-Zertifikat für die Ausfuhr von Arzneimitteln zur Anwendung bei Menschen ohne Zulassung im Ausfuhrland beantragen, Apply for a WHO certificate for the export of medicinal products for human use without a marketing authorization in the exporting country