



Standards for the Management of Cosmetic Sampling Testing (Draft for Comments)

化妆品抽样检验管理规范 (征求意见稿)

National Medical Products Administration

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Translated by ChemLinked

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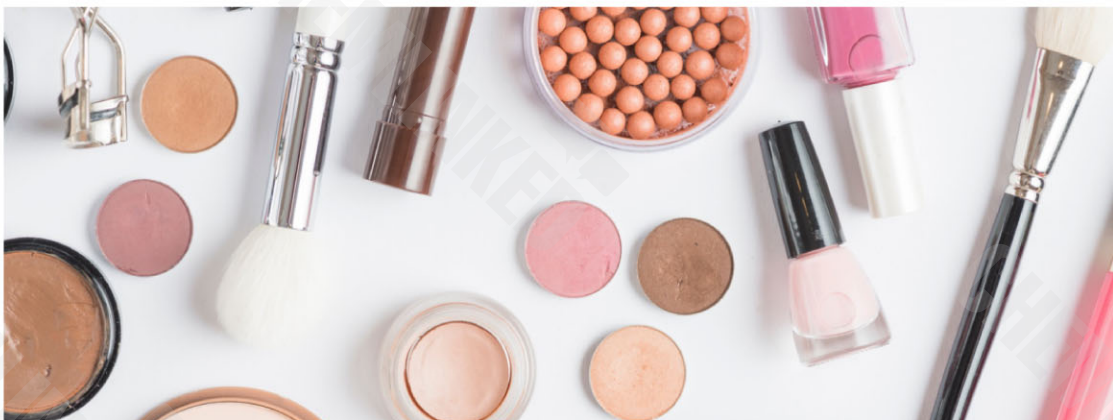
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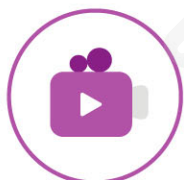


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Chapter I General Rules

Article 1 Legal Basis In order to strengthen the supervision and administration of cosmetics and standardize the sampling testing of cosmetics, this Standard is formulated in accordance with the *Cosmetic Supervision and Administration Regulation*, the *Supervision and Administration Measures on Cosmetics Manufacture and Operation* and other regulations.

Article 2 Scope of Application This Standard shall apply to the cosmetic sampling testing (hereinafter referred to as the sampling testing) involved in the production and operation of cosmetics by the department responsible for medical products supervision and administration within the territory of the People's Republic of China.

Article 3 Management Responsibilities The National Medical Products Administration (hereinafter referred to as the NMPA) is responsible for organizing and carrying out national sampling testing.

The provincial medical products administration department is responsible for organizing and implementing national sampling testing within its administrative area, and is responsible for organizing provincial sampling testing within its administrative area.

The departments responsible for medical products supervision and administration of the people's governments of cities and counties shall implement national and provincial sampling testing in accordance with the arrangement of higher-level departments.

For cosmetics that may have quality and safety risks found in complaint reporting, supervision and inspection, adverse reaction monitoring, and risk monitoring, the department responsible for medical products supervision and administration at or above the county level may conduct special sampling testing.

Article 4 Responsibilities of Testing Institutions National Institutes for Food and Drug Control is responsible for drafting national sampling testing plans and programs, organizing, implementing and providing technical guidance according to the requirements of the NMPA, collecting, analyzing, and reporting the national sampling testing data, and organizing quality analysis and information sharing application.

Cosmetic testing institutions with corresponding testing qualifications shall undertake relevant testing tasks.

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- + Ingredients registration by ICID



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- + Cosmetic test service
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