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## Turkey Updates Application Form and CTD Guide in Relation to the Regulation on Licensing of Medicinal Products for Human Use

## 11 Mar 2022

With respect to the Regulation on Licensing of Medicinal Products for Human Use ('**Regulation**") published in Official Gazette dated 11 December 2021 and numbered 31686, the Turkish Medicines and Medical Devices Agency ("**Agency**") updated the Application Form and CTD (Common Technical Document) Guide.

Certain important matters regarding the updates can be summarized as follows;

- In parallel with the changes introduced with the Regulation, types of hybrid application and related allergen product application have been included.
- Applications regarding processes of prioritized evaluation, conditional licensing (emergency use approval) and exceptional licensing which have been introduced with the Regulation have been regulated.
- A section for medical devices has been included in the application details.
- Regulations regarding the submission of results of preclinical or clinical studies have been included.

You may find the announcement of the Agency regarding the updates through this <u>link</u> (only available in Turkish).

## Related Attorneys

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