

# Regulatory and Policy Updates ANVISA

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- Resolution RDC n° 270/2019 Simplification of the regulatory process for the lowest-risk medical devices.
- Regulatory process for Class I devices (including IVDs) change from cadastro (simplified approval) to a simple notification.
- Effective on 05/02/2019.



- Public Consultation n° 546/2018 Regulations for custom-made devices
- Core elements of the Public Consultation:
- Device manufacturers and importers must be fully licensed by ANVISA;
- Manufacturers of Class III and IV devices must have valid Brazilian Good Manufacturing Practice (BGMP) certifications



- Public Consultation n° 584/2018; 584/2018
   and 586/2018 Updates requirements for Reprocessing & Reuse of Medical Devices.
- Requirements for labeling and for good practices for the processing Medical Devices;
- Anvisa's goal with the proposal of a new RDC is to improve the management of risks associated with the processing of medical devices.



Public Consultations

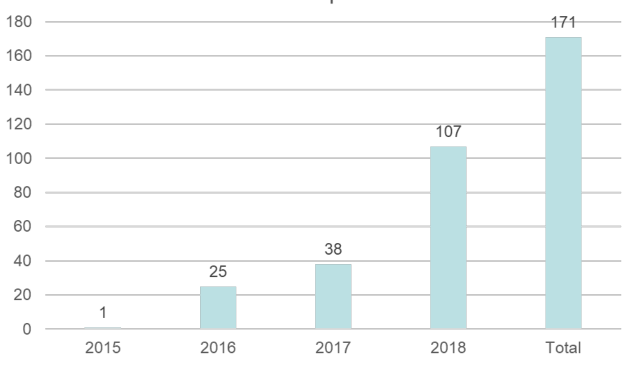
#### How to submit your contribution?

http://portal.anvisa.gov.br/consultas-publicas#/

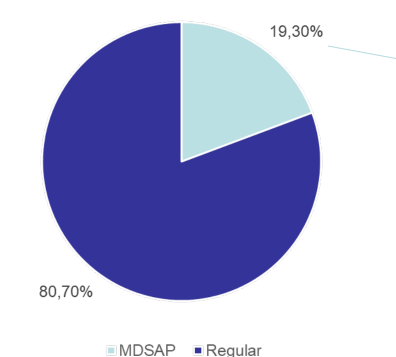
You can also upload documents, such as position papers and send it to ANVISA.



## ANVISA's GMP Certification using MDSAP audit reports



#### Anvisa's GMP Certification 2018



MDSAP Certification 2019

Increasement Projection:
30% to 40%
of the total GMP Certification
to be issued by Anvisa.



### Thank you!

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