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*MDPlaybook 2018: Vesna Janic  
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13485:2012. The revised ISO  
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9001:2008 and not ISO  
9001:2015. This misalignment

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Introducing the new ISO



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device industry, with over  
27,000 certificates  
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?In Europe, ISO 13485  
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the de facto standard for  
the medical device industry.  
?Addresses most or all of

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the quality system

requirements in markets  
including Europe, Australia,  
Japan, Canada, South Korea  
and Brazil, etc.

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ISO 13485, Medical devices – Quality management systems – Requirements for regulatory purposes, is the International Standard for quality management systems for the medical devices sector. Published in 2016,

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it is designed to work with other management systems in a way that is efficient and transparent.

~~ISO - FDA plans to use ISO  
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This standard supersedes earlier documents such as EN 46001 (1993 and 1996) and EN 46002 (1996), the previously published ISO 13485 (1996 and 2003), and ISO 13488 (also 1996). The current ISO 13485 edition was published

# Download File PDF Iso 13485 2016 Standard Published By Iso Group on 1 March 2016.

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The International  
Organization for  
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13485 medical devices

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quality management systems  
standard on March 1, 2016.  
ISO 13485:2016 can be used  
by organizations involved in  
the production, post-  
production, storage,  
distribution, installation,  
servicing, final

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decommission and disposal of  
medical devices.

~~ISO 13485 Medical Devices +  
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On March 1, 2016 the  
International Organization  
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Published the new edition of the ISO 13485 standard. Previously updated in 2003, the revision places more emphasis on the quality management system throughout the supply chain and product lifecycle, as well as on

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device usability and  
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requirements.

~~NEW ISO 13485:2016 GUIDANCE  
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a Final Draft International  
Standard (FDIS)/ISO 13485 on  
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29th of October 2015 for balloting by ISO member countries. The revised standard ISO 13485:2016 was published on 1st March 2016. Summary of the key changes  
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in a number of important  
areas. The following  
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these ...

~~ISO 13485:2016 Revision  
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insiders, ISO pushed ahead with publication of its revised medical device quality management system standard, ISO 13485:2016, despite some controversy that many thought would cause ISO to delay its

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ISO 13485:2016, published by the International Organization for Standardization. Attention is drawn to the following:  
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