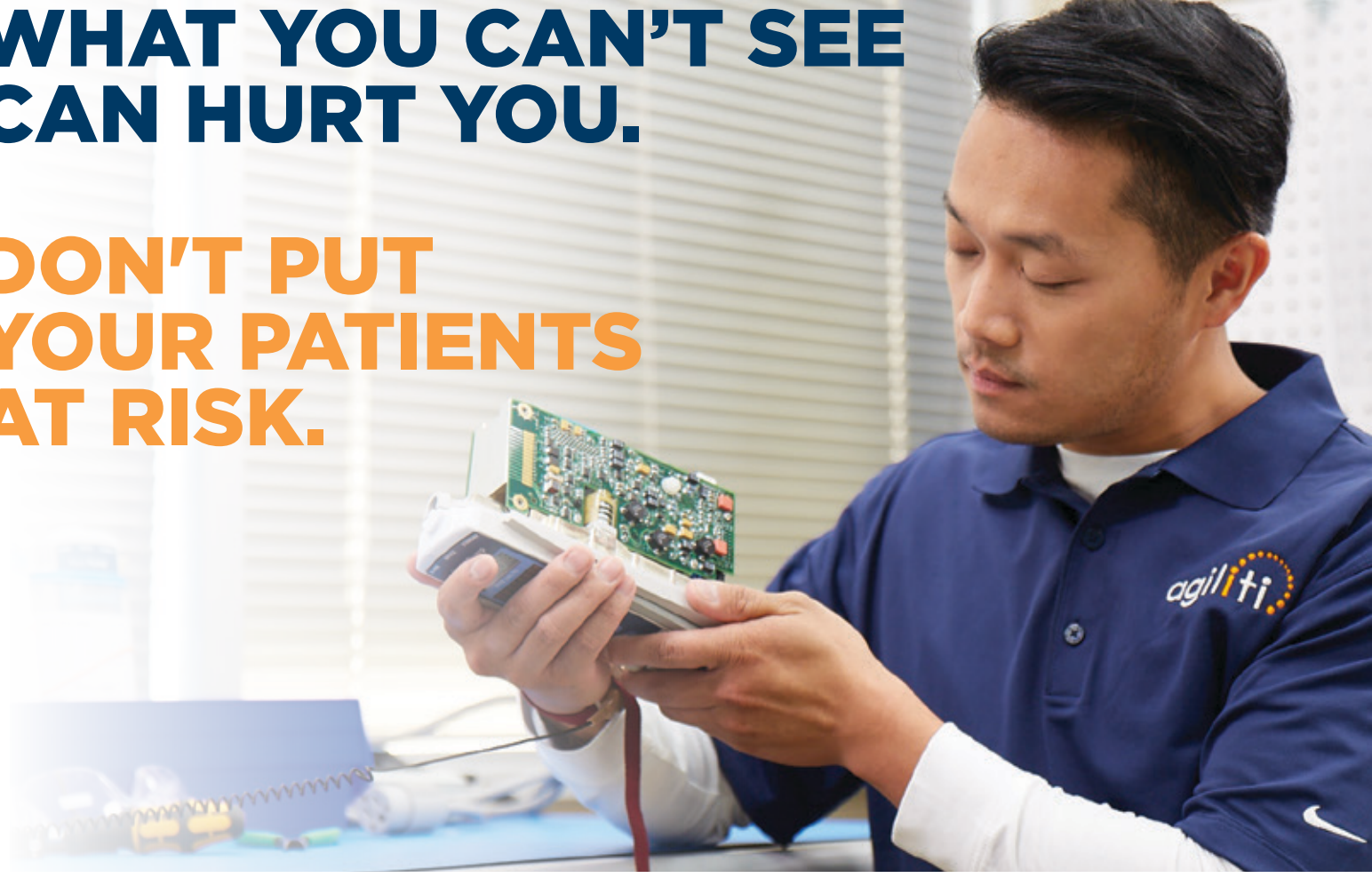


# WHY ISO 13485:2016 IS THE ONLY CHOICE WHEN IT COMES TO PATIENT SAFETY

**WHAT YOU CAN'T SEE CAN HURT YOU.**

**DON'T PUT YOUR PATIENTS AT RISK.**



## ARE YOUR VENDORS UP TO DATE?

The medical device industry has shifted to a higher standard, introducing a growing divide in quality and patient safety assurance.

<p>General <b>ISO 9001:2015</b></p> <p>A generic framework to enhance customer satisfaction in a wide range of industries; not focused on the safety and efficacy of medical devices.</p> <ul style="list-style-type: none"> <li> <b>LANDSCAPING</b></li> <li> <b>FLORAL</b></li> <li> <b>LOGISTICS</b></li> <li> <b>CONSTRUCTION</b></li> <li> <b>PROFESSIONAL CLEANING</b></li> </ul> <p>One is for: <b>REPAIRING LAWNMOWERS</b></p>	VS	<p>Medical Devices <b>ISO 13485:2016</b></p> <p>A rigorous, patient-focused and risk-based quality management system to ensure those in the high-risk medical device industry are delivering the highest level of patient safety and quality.</p> <ul style="list-style-type: none"> <li> <b>MEDICAL DEVICE DESIGNERS</b></li> <li> <b>MEDICAL DEVICE MANUFACTURERS</b></li> <li> <b>MEDICAL DEVICE SERVICERS</b></li> </ul> <p>The other is for: <b>SERVICING VENTILATORS</b></p>
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## THERE IS NO SUCH THING AS A HARMLESS SHORTCUT

Third-party ISO 13485:2016 certification ensures patient safety isn't compromised at any step of the medical device lifecycle.

### 3 KEY REQUIREMENTS (Out of 80+) to illustrate why ISO 13485:2016 is the only acceptable standard.

RISK-BASED APPROACH	CUSTOMER FEEDBACK	RECALL MANAGEMENT
<p><b>WHY IT MATTERS:</b></p> <p>New standards require a risk-based approach at every step of a provider's Quality Management System (QMS), with consideration for the impact of medical device technology — from software to documentation.</p>	<p><b>WHY IT MATTERS:</b></p> <p>ISO 9001 focuses on customer satisfaction; ISO 13485 focuses on customer feedback on device performance. The difference? Added focus on keeping patients safe, not just happy.</p>	<p><b>WHY IT MATTERS:</b></p> <p>A rigorous recall management process to ensure quick customer notification of recalls — active identification of impacted devices. Minimizes both patient risk and hospital disruptions caused by recalls.</p>

**YOUR DOCTORS AND NURSES WON'T FAIL HER.**

**MAKE SURE YOUR EQUIPMENT DOESN'T EITHER.**



## THE CHOICE IS CLEAR

ISO 9001:2015 vs ISO 13485:2016

	ISO 9001	ISO 13485
Determine, monitor and review external and internal issues	✔	✔
Use risk-based thinking for planning and management	✔	✔
Provide and maintain resource monitoring	✔	✔
Monitor and analyze implementation with an internal audit program	✔	✔
Utilize risk-based approaches that consider patient safety through every step		✔
Provide early warning of quality issues through a feedback system		✔
Investigate equipment incidents reported through customer feedback		✔
Maintain the suitability and effectiveness of the quality management system		✔
Control medical devices with regard to advisory notices (product recalls, etc.)		✔
Control the work environment to prevent contamination		✔
Retain obsolete controlled documents for the life of the medical device		✔
Staff trained and supervised to work in special conditions (bloodborne pathogens, etc.)		✔
Control of customer property, including confidential health information		✔
Validation of computer software		✔
Full traceability of devices, including documents and records		✔

Align your patient safety goals with a vendor committed to the highest standards in the country.

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