CASE FOR QUALITY
Safe Space / Medical Device Information Analysis & Sharing (MDIAS)

During the medical device development process and in real-world environments, an abundance of data is collected. Currently siloed within medical device companies for exclusive organizational use and for patient privacy, these data sets vary in quality and format. Developing a manner to collect and analyze data would help identify trends in device quality and safety to provide powerful insights for improving outcomes while safeguarding data.

The Safe Space/ MDIAS initiative focuses on a data-sharing collaborations to analyze and share medical device data from various public and non-public sources to improve healthcare outcomes. Data is safeguarded by an independent trusted third party to foster broad participation and engagement.

How To Get Involved
Learn more and Share your ideas
Get to know Safe Space / MDIAS! If you have an idea for this project, we want to hear from you.

For further information contact the MDIC Case for Quality Team at CfQcc@mdic.org.

FDA does not lead Collaborative Communities, and their work does not replace established regulatory mechanisms. However, FDA may choose to participate in a Collaborative Community and support, leverage, and/or adopt solutions that emerge.

ABOUT MDIC
The Medical Device Innovation Consortium (MDIC) is a public-private partnership that brings together representatives of the FDA, NIH, CMS, NIST, and other agencies, industry, non-profits, and patient advocacy organizations to improve the processes for development, assessment, and review of new medical technologies.

MDIC coordinates the development of methods, tools, and resources used in managing the total product life cycle of a medical device to improve patient access to cutting-edge medical technology.

We are driving faster, safer, and more cost-effective innovation for patient benefit.

CASE FOR QUALITY COLLABORATIVE COMMUNITY (CfQcc)
The Safe Space / MDIAS Initiative is an effort of the Case for Quality Collaborative Community and offers a unique forum for medical device stakeholders to move beyond baseline regulatory compliance activities and collaboratively develop sustained predictive practices. These predictive practices could advance medical device quality and safety to achieve better patient outcomes.

Collaborative Communities provide a forum for public and private sector members to proactively work together to achieve common objectives and outcomes.

LEARN MORE ABOUT MDIC’S CfQcc
Learn more about MDIC’s CfQcc by visiting www.mdic.org.