

	5		
QT9 QMS + Your Logo		Sam Sturm Corp Corp Logout	User To Do
Global Manage ISO Functions Doc. Control Products Reporting	Training Projects / Tasks Maintenance	Safety Customers Suppliers Help	
Corrective Actions	ail CAR	E	lack To List
Corrective Action Type		Priority / Assignment	., ,
CAR Type Customer Complaint Problem Type -= Select Problem =-	~	Priority Critical ~	
		Resp. Party: Christian Reyes	QT9QM
Cecception Material Info Additional Data Root Cluste User Defined Analysis Related Files	Sup. Response Tasks Verification Approves	i / Mgmt. Notes Timeline Related Files	Sign In
ICS Demo Files	Supplier Uploaded Files		
Add file	There are currently no files attached.		Password
			Remember me Forget Passwor
There are currently no files attached.			
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There are currently to thes attached.			Sign In
Level on Recognitive Statistics Send to Recognitive Statistics (B) Salest for Approximate Statistics (B) Salest for Approximate Statistics (B) Salest Statis	Save CAR	Drillete CAR	Sign In

Paper Required

Modules

Traceability

0%

23+

100%

Quality Management Software Made Easy

Automate Regulatory Compliance

Life Sciences | Aerospace | Automotive | Manufacturing | Foods & Beverages

FDA Compliance

- 21 CFR Part 11
 Electronic Signatures
- 21 CFR Part 210/211
 Pharmaceutical Companies
- V 21 CFR Part 820

 Medical Device Manufacturers
- FDA 510 K

 Medical Device Premarket

ISO Compliance

- ISO 9001
 Quality Management System
- ISO 13485

 Medical Device Makers
- ♣ ISO 14001 Environmental Management
- Laboratory Testing & Calibration

Standards

- AS9100
 Aerospace & Defense Industry
- A HACCP
 Food Safety Industry
- Automotive Industry
- SQF Food Safety
 Food Safety Managment

Unify Quality Management Processes

23+ Modules Included | Connect Quality Processes In One Unified Suite

Core Modules

a Audit Management

Corrective Actions (CAPA)

Preventive Actions (CAPA)

Management Review

Nonconforming Products

A Risk Assessments

Customer Modules

Customer Feedback

Customer Complaints

Customer Surveys

Customer Web Portal

Document Modules

Document Control

Link to Training

Revision Control

₩ Master List of Records

Obsolete Docs/Legacy Docs

Document Viewing Portal

Product Modules

Deviation Management

≠ ECR/ECN

■ FMEA

Inspections

A Material Review Board

Supplier Modules

Supplier Evaluations

Supplier Surveys

Supplier Web Portal

Management Modules

E Change Control

Employee Training

Project Management

Safety Incidents

Maintenance Modules

Calibrations

Preventive Maintenance

Unlock Insights with Real-Time Data

Personalized Super-Grids

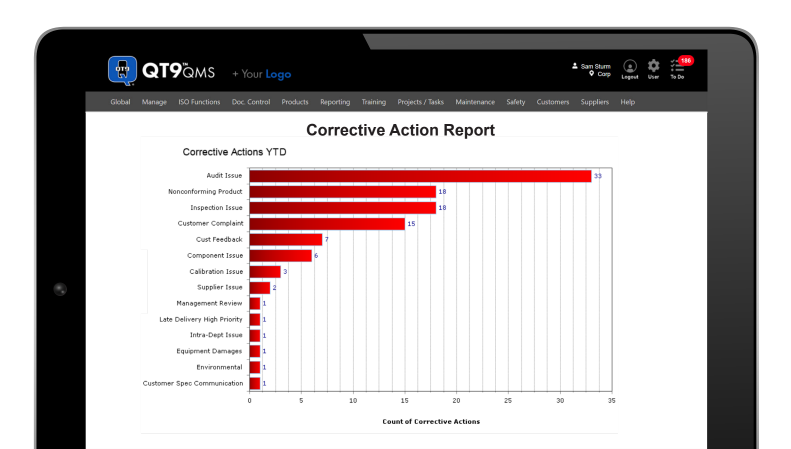
Create multiple views of your data & save each view for future reference. Filter, sort & group data the way you want to see it.

Digitally Transform Quality

Check the status of items 24/7. Enable forward-thinking intelligence to take quality to the next level.

User-Friendly Web Portals

Centralize data & increase visibility from your suppliers, customers & employees with QT9's easy-to-use web portals.



The Ultimate QMS Platform



Flexible Platform

Expand your capabilities with a 100% Cloud-Based Web Subscription,
On-Premise Web Purchase or
Windows Client Desktop Cloud.



Highly Scalable Solution

Get unlimited scalability and dynamically populate forms with data inherited from one process to the next with QT9's extensible & modular architecture created for growth.



Total Traceability

Make informed decisions with date/
time/name stamped entries to see
which user made the changes and when
the changes were made.



Unlimited Training

No matter who is working at your company in quality, QT9 includes unlimited web training and support for all employees.



Free Upgrades

Future-proof your organization with access to the latest software features.

Ensure your software is always current with QT9™ QMS.



Modern Technology

From fast-growing start-ups to global enterprises, QT9 gives you the tools to automate the evolving quality and compliance requirements as you grow.

Start a Free Trial • Visit QT9QMS.com



The QT9 Experience

- User-Friendly Design
- User To-Do List
- **Unlimited Support**
- Online Videos
- □ Webinars
- ¶ Knowledge-Base
- Unlimited Scalability
- **Proven Reliability**

Digitally Transform Quality

■ Web-Based Platform

Multi-Site Architecture

Interconnected Modules

Email Alerts

Approval Management

Priority Management

Centralize Related Files

⇄ Import/Export Tools

eSignatures

User Rights Access

+ Custom Data Fields

🚹 Unlimited Data

Ready-To-Go

Access and process items virtually anywhere.

Centralize records across multiple sites.

Centralize documents and records.

Send users reminders with hotlinks.

Assign approvers based on procedures.

Set priority levels based on urgency

Attach unlimited related files to action items.

Transfer data between systems.

FDA 21 CFR Part 11 electronic approvals.

Designate what users are allowed to see.

Track your unique data-driven analytics.

Include unlimited file attachments.

Get going with ready-made templates.

^{*} Requires an active QT9 QMS subscription.