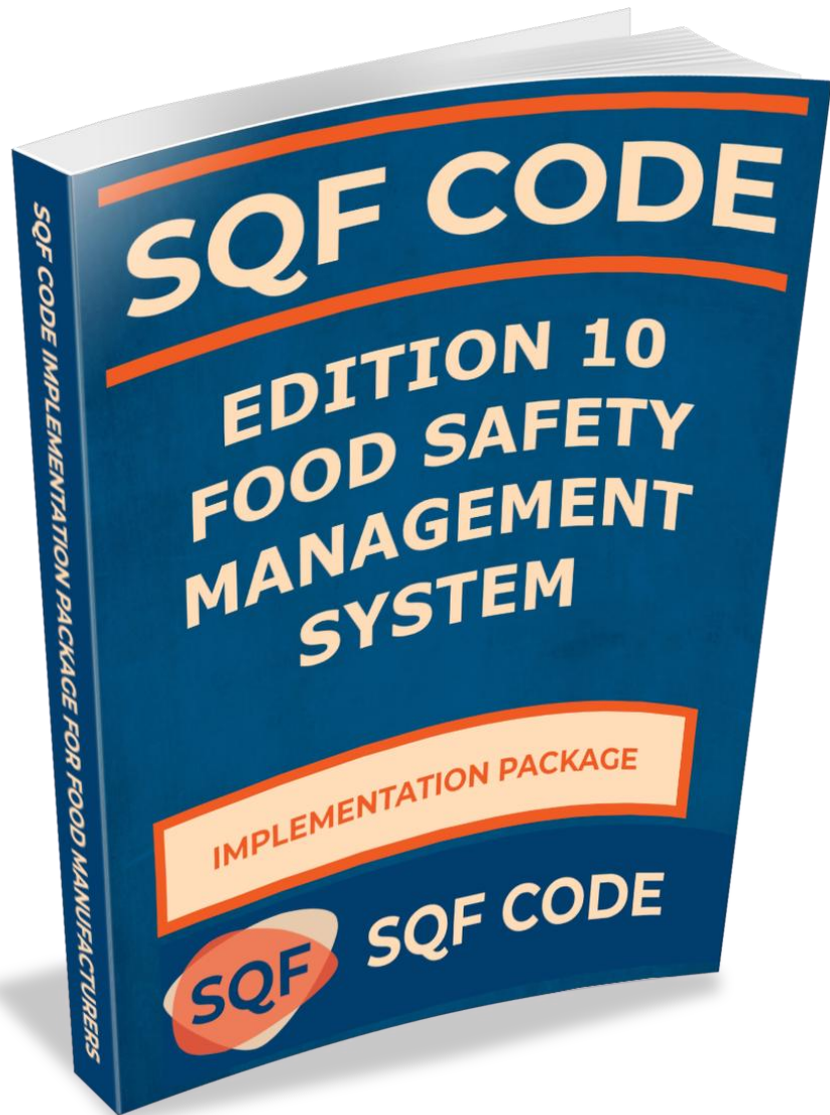


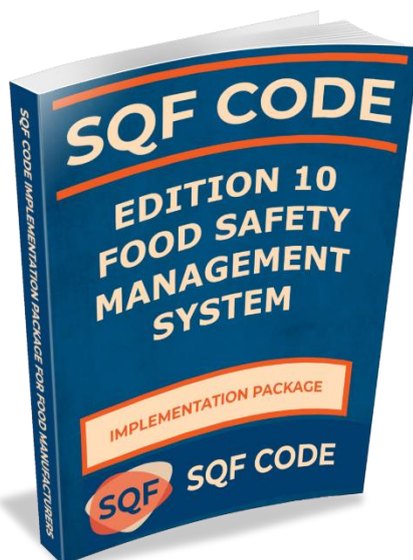
New 2026 Implementation Package Compliant with SQF Food Safety Code: Food Manufacturing Edition 10 and the latest CODEX HACCP Guidelines



This is an ideal package for Food Manufacturers looking to achieve certification to the SQF Food Safety Code: Food Manufacturing Edition 10

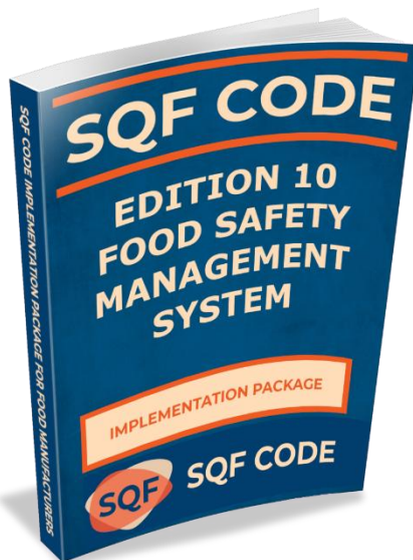


The SQF Food Safety Management System Implementation Package includes a combination of comprehensive documentation, guidance, implementation tools and training.



Included in the SQF Food Safety Management System Implementation Package:

- ✓ Comprehensive Procedures Manual
- ✓ Supplementary HACCP Tools & Documents containing the HACCP Calculator
- ✓ Laboratory Quality Manual
- ✓ Training Modules
- ✓ FSQMS, Verification and Validation Record Templates
- ✓ Free online support via e-mail
- ✓ Allergen Management Module & Risk Assessment Tool
- ✓ Supplier Risk Assessment Tool
- ✓ Product Development Module
- ✓ Complaint Management Guidelines & Analyser
- ✓ Internal Audit Schedule Risk Assessment Tool and Template
- ✓ Food Fraud Risk Assessment Tool
- ✓ Food Defence Assessment Tool
- ✓ Implementation Workbook
- ✓ User guide



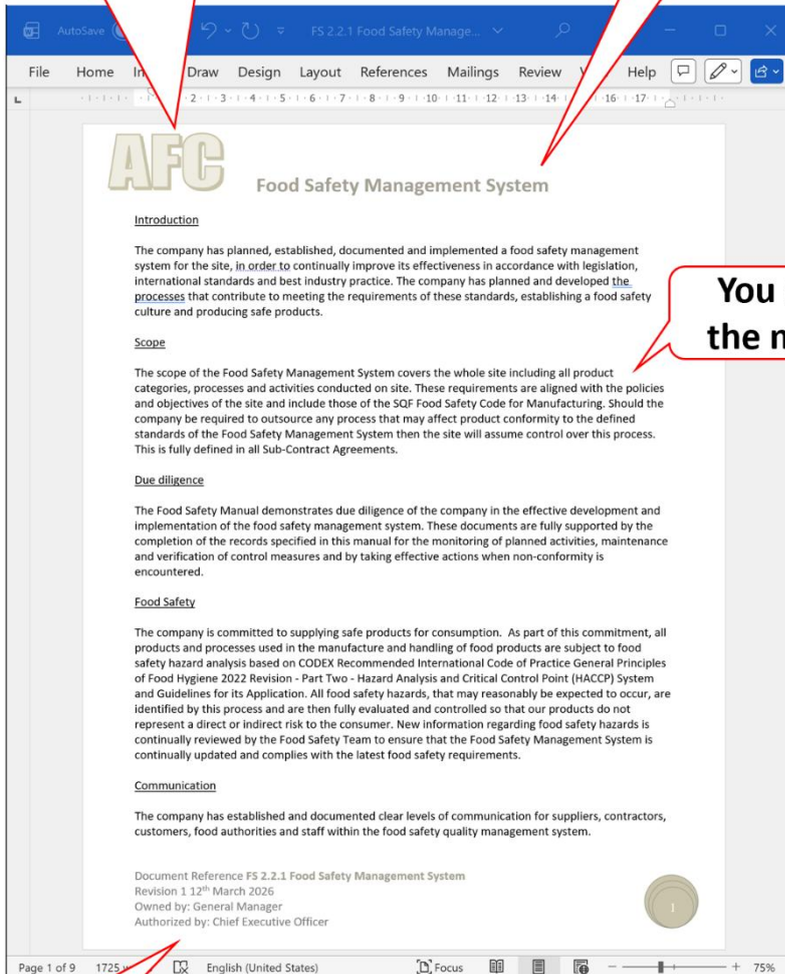
[To order the SQF Edition 10 Food Safety Management System Implementation Package click here](#)

Food Safety Management System & Prerequisite Programme Procedures

The main documents are provided in Microsoft Word format and are easily edited to suit your organisation.

For example put your company logo or name and address in the header

You can edit the header



You can edit the main text

You can edit the footer

The package contains a comprehensive set of editable Food Safety Management System & Prerequisite Procedures written in Microsoft Word (US English) format that match the clauses of the SQF Food Safety Code: Food Manufacturing Edition 10.

These are the procedure templates for Sections

2.1 Management Commitment

2.2 Document Control and Records

2.3 Specifications, Product Development, and Supplier Approval

2.4 Food Safety System

- FS 2.1 Management Commitment
 - FS 2.1.1.1 Food Safety Policy
 - FS 2.1.1.2 Food Safety Culture
 - FS 2.1.1.2 Food Safety Culture - Expected Behaviors
 - FS 2.1.1.3 Food Safety Culture Planning Matrix
 - FS 2.1.1.4 Food Safety Objectives
 - FS 2.1.1.4 Responsibility and Authority
 - FS 2.1.1.4 Appendix Organizational Chart
 - FS 2.1.1.4 Appendix Job Descriptions
 - FS 2.1.2 Management Review
 - FS 2.1.3 Complaint Management
 - FS 2.1.3 Note - How to reduce your Complaint levels
 - FS 2.1.3A Annual Complaints Analyzer
 - FS 2.1.3B Annual Complaints Analyzer Instruction
- FS 2.2.1 Food Safety Management System
 - FS 2.2.2 Document Control
 - FS 2.2.3 Records
- FS 2.3.1 Product Development
 - FS 2.3.1A Development Supplementary Documents
 - FS 2.3.2 Specifications
 - FS 2.3.2A Material Acceptance Record
 - FS 2.3.3 Contract Manufacturers
 - FS 2.3.4 Approved Supplier Program
 - FS 2.3.4A Supplier & Material Risk Assessment
- FS 2.4.1 Food Legislation
- FS 2.4.2 Good Manufacturing Practices
 - FS 2.4.3 Food Safety Plan
 - FS 2.4.3A Additional HACCP Tools
- FS 2.4.4 Product Sampling, Inspection and Analysis
 - FS 2.4.4A Laboratory Quality Manual
 - FS 2.4.4B Product Sampling Supplementary Documents
- FS 2.4.5 Non-conforming Materials and Product
 - FS 2.4.6 Product Rework
 - FS 2.4.7 Product Release
- FS 2.4.8 Environmental Monitoring
 - FS 2.4.8A Appendix Environmental Monitoring

PACKAGE DOCUMENT EXAMPLES

AFC Management Commitment

Introduction

Senior management demonstrate clear and visible commitment to the food safety management system by establishing and implementing, then fully communicating and supporting its policies, procedures and objectives. Senior Management is committed to continually improve the effectiveness of the food safety management system by regular audit, review and proactive actions.

Scope

The scope of the Food Safety Management System includes all products manufactured on site and activities conducted on site.

The scope is aligned with the policies and objectives of the site and includes the commitment to fully meet the requirements of the SQF Food Safety Code: Food Manufacturing, Edition 9.

Procedure

The Senior Management has a total commitment to food safety, observing all legal, moral and ethical codes and this is the concern of every employee.

Senior management demonstrate clear and visible commitment by:

- Establishing and implementing a Food Safety Policy,
- Communicating and Maintaining the Food Safety Policy,
- Establishing and Implementing Food Safety Objectives,
- Communicating and Maintaining the Food Safety Objectives
- Leading and supporting a food safety culture within the site
- Conducting regular proactive management reviews and communicating outputs,
- Communicating commitment to satisfying customer requirements including food safety, quality and service
- Supporting and planning the development and operation of the Food Safety Management systems.
- Ensuring the food safety management system is maintained when changes are planned and implemented.
- Establishing documentation required for the effective development, implementation and updating of the food safety management system and communicating pertinent information throughout the organization.
- Providing the human and financial resources, and training, to manage the Policies and Objectives effectively.
- Providing the infrastructure and work environment to manage the Policies and Objectives effectively.
- Promoting an ethic of continuous improvement throughout the company.

Document Reference FS 2.1 Management Commitment
Revision 0 1st August 2023
Owned by: Quality Manager
Authorized by: Managing Director

Expected Behaviors of all Personnel



- ✓ Contribute to company objectives
- ✓ Compliance with company procedures
- ✓ Correctly completing documentation and records as required by your role within the organisation
- ✓ Adhere to Hygiene rules and comply with expected personnel standards
- ✓ Report non-conforming products or equipment
- ✓ Report any issues or areas of concern that may affect product safety, authenticity, legality or quality
- ✓ Report any problems with pests
- ✓ Ensure site security procedures are followed and unknown visitors are challenged
- ✓ Adopt a 'clean as you go' policy
- ✓ Contribute to hygiene and housekeeping standards
- ✓ Make suggestions for improvement

Environmental Monitoring Priorities

Open product areas:
High risk (chilled and frozen)
High care (chilled and frozen)
Ambient high care
Low risk
Flow & entrances to the above areas

Enclosed product areas:
Warehouses
Storerooms
Flow & entrances to the above areas

Non-product areas:
Canteens
Laundries
Offices
Flow & entrances to the above areas

Priority Order for Environmental Sampling

Environmental Monitoring Schedule

Standards for Plant and Equipment may need to be established during production runs as well based on product conformance throughout a production run.

Surface	Frequency	Target	Level	Target	Level
Food Contact Surface – Filler Nozzle	Weekly	TVC	Monthly	Salmonella	Absent
Food Contact Surface – Foil Lidding	Weekly	Y&M < 100	Monthly	Listeria	Absent
Non-Food Contact Surface – Inside Door Filler Cabinet	Weekly	Enterococci < 10	Monthly	E.Coli O157	Absent
Non-Food Contact Surface – Cleaning Equipment	Weekly	E.Coli < 1	Monthly	Staph aureus**	Absent
Non-Food Contact Surface – Floor under Filler	Weekly	TBC	Quarterly	Contact	< 10
Non-Food Contact Surface – Outside Storage Tank	Monthly	TBC	Quarterly	Non-contact	TBC
Non-Food Contact Surface – Drain	Monthly	TBC	Quarterly	Non-contact	TBC
Non-Food Contact Surface – Wall	Monthly	TBC	Quarterly	Non-contact	TBC
Non-Food Contact Surface – Floor near Entrance	Monthly	TBC	Quarterly	Non-contact	TBC
Non-Food Contact Surface – Hand Wash Sink	Monthly	TBC	Quarterly	Non-contact	TBC

AFC Food Safety Culture

Introduction

The company recognizes that a successful food safety culture is the product of individual and group values, attitudes, competencies and patterns of behavior that determine the commitment to, and the style and proficiency of the food safety management system. The site's senior management plan for the development and continuing improvement of food safety culture.

Senior management are responsible for delivering a "It is how we do things here" food safety culture by:

- Leadership – starting from the top
- Demonstrating visible commitment
- Effective communication of company philosophy and policy
- Ensuring there is accountability from the top of the organization to the bottom
- Developing employee confidence and mutual trust
- Developing reward schemes including 'Employee of the Month' award
- Ensuring all employees are accountable, engaged and understand the value of integrity and proactivity
- Developing an action plan for the development and continuing improvement of food safety culture

Developing a Food Safety Culture

A successful food safety culture can be achieved only by following safe working practices and procedures developed through effective hazard analysis, training and experience. In order to achieve these aims, a robust Food Safety Hazard Analysis Critical Control Points System (HACCP) has been introduced following a full hazard analysis of all food related operations. All instructions and control mechanisms within the Food Safety (HACCP) System are designed to control any risk to food safety.

To ensure success of this policy Senior Management are directly responsible for food safety by ensuring adequate; organization and support, equipment and facilities, training and education of all employees, reviewing and auditing performance, and driving continuous improvement. Detailed organizational arrangements and food safety responsibilities for all levels of management are contained in the food safety and manual and job descriptions.

Achievement of this policy involves all staff being individually responsible for the quality of their work, resulting in a continual improvement culture and working environment for all. All employees are provided with the food safety training necessary to enable them to perform their tasks and are responsible for ensuring that they do so in a hygienic manner so that the safety of the food they handle is not put at risk. All employees are required to co-operate with any authorized person to ensure that customer, statutory and regulatory obligations are properly complied with. Employees are encouraged and required to notify management about actual or potential food safety issues and are empowered to act to resolve food safety issues within their scope of work.

Document Reference FS 2.1.1.2 Food Safety Culture
Revision 0 1st August 2023
Owned by: Quality Manager
Authorized by: Managing Director

AFC Approved Supplier Program

- ✓ Microbiological contamination
- ✓ Chemical contamination
- ✓ Physical contamination
- ✓ Allergens and possible allergen contamination
- ✓ Possible substitution or fraud
- ✓ Effect on product quality

Consideration is given to the significance of a material to the quality of the final product. The results of the risk analysis dictate the criteria for supplier assurance, testing and acceptance of raw materials and procedures for supplier monitoring. All risk assessments are reviewed when there are changes to materials and at a minimum annually.

Supplier Category Rating
Final Ingredients/Contract Packager
Raw Ingredient/High Risk Service
Contact Packaging
Non-Contact Packaging
Low Risk Service

Document Reference FS 2.3.4 Approved Supplier Program
Revision 0 1st August 2023
Owned by: Quality Manager
Authorized by: General Manager

AFC Food Safety Plans

Introduction

We are a leading food company committed to producing safe and legal products in line with legislation and to continuously improve our standards of hygiene, quality and safety in relation to both our product range and the environment in which we manufacture these products. As part of this commitment, all products and processes used in the manufacture of food products are subject to hazard analysis based on the Codex Alimentarius HACCP principles and the requirements of SQF Food Safety Code: Food Manufacturing.

The Food Safety Manual demonstrates due diligence of the company in the effective planning, development and implementation of the food safety management system. These documents are fully supported by the completion of a Food Safety plan and the records specified in this manual for the monitoring of planned activities, maintenance and verification of control measures and by taking effective actions when non-conformity is encountered. All food safety hazards, that may reasonably be expected to occur, are identified by this process and are then fully evaluated and controlled so that our products do not represent a direct or indirect risk to the consumer.

The Food Safety Management System is fully supported by established verification procedures and validation of the control measures/combination of control measures that are implemented through good manufacturing practices (when applicable) or the Food Safety plan.

HACCP Application

The Company Food Safety System has been developed based on CODEX Recommended International Code of Practice General Principles of Food Hygiene 2022 Edition - HAZARD ANALYSIS AND CRITICAL CONTROL POINT (HACCP) SYSTEM AND GUIDELINES FOR ITS APPLICATION - SECTION 19: APPLICATION

- 19.1 Assemble HACCP Team and Identify Scope (Step 1)
- 19.2 Describe product (Step 2)
- 19.3 Identify intended use and users (Step 3)
- 19.4 Construct flow diagram (Step 4)
- 19.5 On-site confirmation of flow diagram (Step 5)
- 19.6 List all potential hazards that are likely to occur and associated with each step, conduct a hazard analysis to identify the significant hazards, and consider any measures to control identified hazards (Step 6/ Principle 1)
- 19.7 Determine the Critical Control Points (Step 7/ Principle 2)
- 19.8 Establish validated critical limits for each CCP (Step 8/ Principle 3)
- 19.9 Establish a Monitoring System for Each CCP (Step 9/ Principle 4)
- 19.10 Establish corrective actions (Step 10/ Principle 5)
- 19.11 Validation of the HACCP Plan and Verification Procedures (Step 11/ Principle 6)
- 19.12 Validation of the HACCP Plan
- 19.12.1 Verification Procedures

Document Reference FS 2.4.3 Food Safety Plans
Revision 0 1st August 2023
Owned by: Quality Manager
Authorized by: General Manager

PACKAGE DOCUMENT EXAMPLES

AFC Internal Audits

Introduction

The company has established, documented and implemented an internal audit system to verify the Food Safety Management System is effective and continually improved.

Scope

The scope of the Internal Audit System covers all aspects of the food safety management system includes all products manufactured on site, activities conducted on site and all applicable requirements of the SQF Food Safety Code for Manufacturing.

Procedure

The Senior Management has a total commitment to the food safety management system and provides adequate resource in the form of trained and qualified personnel to carry out a comprehensive Internal Audit Schedule. Internal audits are performed to confirm that company management systems are working effectively and to promote continuous improvement. The company philosophy is to audit, review and improve.

The Internal Audit Schedule is planned annually and is designed to comprehensively cover all areas of the Food Safety Management system including procedures, policies and activities. All areas are audited in full at least once per annum; the frequency of audit of each area is based on risk assessment by the Quality Manager.

Document Reference FS 2.5.4 Internal Audits
Revision 0 11th March 2026
Owned by: Quality Manager
Authorized by: General Manager

AFC Internal Audits

Internal Audit Procedure

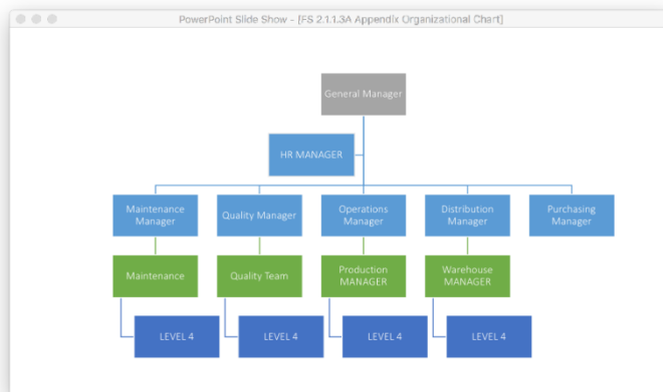
Internal auditors are required to be trained and competent in internal audit procedures, and where practical independent of the function being audited.

Internal Auditors are responsible for carrying out the procedure as described below:

- The site internal audit schedule determines which audits are to be carried out. Each audit is assigned to an auditor independent of that area whenever possible. The auditor must make sure they have the correct audit checklist form to carry out the audits.
- A date and time for the audit to take place must be agreed with the department. A representative from the department must be present during the audit.
- The auditor uses a specific audit form and checklist designed by the Quality Manager for each department or area. Objective evidence is recorded to verify compliance and/or non-compliance.
- The audit report is rated based on the following criteria:
 - RED** – Critical/Major Non-conformance(s) identified and imminent risk. Immediate documented Corrective Action is required and a written follow-up necessary.
 - AMBER** – Minor Non-Conformance(s) identified there is a potential risk. The Corrective Action required is documented and a verbal follow up is required.
 - GREEN** – Satisfactory or Positive with comments or suggestions for improvement
- When the audit is completed and the report given a rating. Positive as well as negative comments are included in the report. Major Non-conformities are immediate highlighted to the department manager, who is responsible for the corrective and preventive action without undue delay.
- The Department Manager reviews the audit findings with the auditor and agrees timescales to complete corrective action for the major and minor non-conformances.
- The Department Manager then signs and retains a copy of the report which includes details of the non-conformances, proposed corrective actions and the agreed time scale to complete the corrective actions. If the audit rating is red then an immediate corrective action plan is reported to the Quality Manager.
- The Departmental Manager is responsible for documenting the corrective actions taken for all the non-conformances raised.

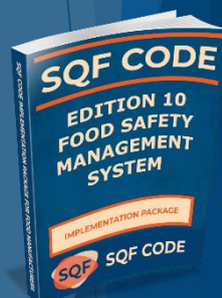
Document Reference FS 2.5.4 Internal Audits
Revision 0 11th March 2026
Owned by: Quality Manager
Authorized by: General Manager

	A	B	C	D	E	F	G	H	I	J	K	L	M	N
1	SQF Internal Audit Schedule													
2														
3	High Risk - Quarterly													
4	Medium Risk - Twice per Year													
5	Low Risk - Annually													
6														
7	Procedure to be covered by Internal Audit													
8	FS 2.1.1 Management Responsibility	Low												
9	FS 2.1.2 Management Review	Low												
10	FS 2.1.3 Complaint Management	Medium												
11	FS 2.2.1 Food Safety Management System	Low												
12	FS 2.2.2 Document Control	Low												
13	FS 2.2.3 Records	Low												
14	FS 2.3.1 Product Formulation and Realization	Medium												
15	FS 2.3.2 Specifications (Raw Material, Packaging, Finished Product and Serv	Medium												
16	FS 2.3.3 Contract Manufacturers	Medium												
17	FS 2.3.4 Approved Supplier Program	High												
18	FS 2.4.1 Food Legislation	Medium												
19	FS 2.4.2 Good Manufacturing Practices	Medium												
20	FS 2.4.3 Food Safety Plan	High												
21	FS 2.4.4 Product Sampling, Inspection and Analysis	High												
22	FS 2.4.5 Non-conforming Materials and Product	High												
23	FS 2.4.6 Product Rework	High												
24	FS 2.4.7 Product Release	Medium												
25	FS 2.4.8 Environmental Monitoring	Medium												



These are the procedure templates for Sections
2.5 SQF System Verification
2.6 Product Traceability and Crisis Management
2.7 Food Defense and Food Fraud
2.8 Allergen Management
2.9 Training

FS 2.5.1 Validation and Effectiveness
FS 2.5.2 Verification Activities
FS 2.5.3 Corrective and Preventative Action
FS 2.5.3A Root Cause Analysis
FS 2.5.3B Corrective Action Request
FS 2.5.3C Preventative Action Request
FS 2.5.4 Internal Audits and Inspections
FS 2.5.4A Audit and Inspection Schedule
FS 2.6.1 Product Identification
FS 2.6.2 Product Trace
FS 2.6.2A Traceability System Diagram
FS 2.6.2B Batch Identification System
FS 2.6.3 Product Withdrawal and Recall
FS 2.6.3A Recall Template
FS 2.6.4 Crisis Management Planning
FS 2.7.1 Food Defense Plan
FS 2.7.1A Food Defense Threat Assessment
FS 2.7.2 Food Fraud
FS 2.7.2A Food Fraud Assessment Template
FS 2.8 Allergen Management
FS 2.8.1A Allergen Management Tool
FS 2.8.1B Allergen Clean Validation
FS 2.8.1C Allergen Clean Verification
FS 2.8.1D Ingredient Allergen Management - Color Coding
FS 2.8.1E Allergens
FS 2.8.1F Allergen Management Records
FS 2.9 Training
FS 2.9A Sample Work Instruction



PACKAGE DOCUMENT EXAMPLES

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Comments Editing Share

AFC Product Trace

Introduction

The company has established, implemented, documented a trace system for all product components. This procedure defines how materials and finished products are uniquely identified and traceable as required by the Food Safety Management System and in compliance with all regulatory requirements in the country of production and sale.

Scope

This procedure applies to all process steps where controls are exerted include raw material intake, ingredients and packaging, work-in-progress, final product and dispatched shipment to customer.

Procedure

A system for identification and traceability of product batches is maintained which, in the event of food safety incidents will enable tracking of material batches (including processing aids) through to distributed batches of finished product using label detail and expiry code. For a traceability to be enacted the product expiry code must be known.

The company traceability system takes both the form of documented records and plc program, which enables a full product history to be produced in a timely manner.

Traceability records by Label and Expiry date are maintained and retained for all product batches. This allows the site to trace materials from goods receipt to customer for every delivery. Records are maintained of raw material and packaging usage and finished product volumes. Procedures ensure that label use is reconciled, and any inconsistencies investigated and resolved. Finished product labels are retained – see FS 2.6.3C Label Retention and Check.

Reworked material also remains identifiable and traceable. Where rework or any reworking operation is performed, traceability is maintained by completing traceability records to the finished product to ensure that product safety or legality is not compromised e.g. allergy status, identity preservation and ingredient declarations.

The traceability will provide details on all parts of the product from raw material intake through to filling time. The traceability entails tracing a product backwards from finished package to its raw materials, ensuring that all associated chemical, physical and microbiological tests, cleaning of equipment and all relevant paperwork has been completed and is within specification.

A mass balance exercise is conducted from of raw material and packaging usage and finished product volumes to ensure that all finished products are accounted for.

Document Reference FS 2.6.2 Product Trace
Revision 0 11th March 2026
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AFC Product Trace

For all products, the following information is traceable from the product expiry code:

Stage	Details	Relevant Record
Raw Material Intake	Materials, Time, Date, Temperature, Batch Code, Supplier, Amount, COC or COA	QMR Raw Material Intake Record
Packaging Intake	Packaging Material, Batch Code, Date, Supplier, Amount, COC or COA	QMR Packaging Intake Record
In-Process batches	Batch Name, Records of all Ingredients mixed including Reworked material. Batch Code	QMR In-Process Record
Process Records	Batch Name, Hot/Cold Temperature and Time. Batch Code	QMR Process Record
Bulk Storage Records	Product, Temperature and Time. Batch Code	QMR Bulk Storage Records
Production Records	Product, Time, Date, Label, Expiry Code, Code of Packaging, Temperature, Quantity, Product & Packaging Reconciliation. Batch Code	QMR Production Records
Storage Record	Product, Time, Date, Label, Expiry Code	QMR Storage Record
Dispatch Records	Product, Time, Date, Label, Expiry Code, Amount, Customer	QMR Dispatch Record
Critical Control Records	For all Control Points	QMR Critical Control Records
Cleaning Records	For all stages	QMR Cleaning Records
Delivery Records	Products, Customer & Location Time, Date, Label, Expiry Code, Amount	QMR Delivery Record

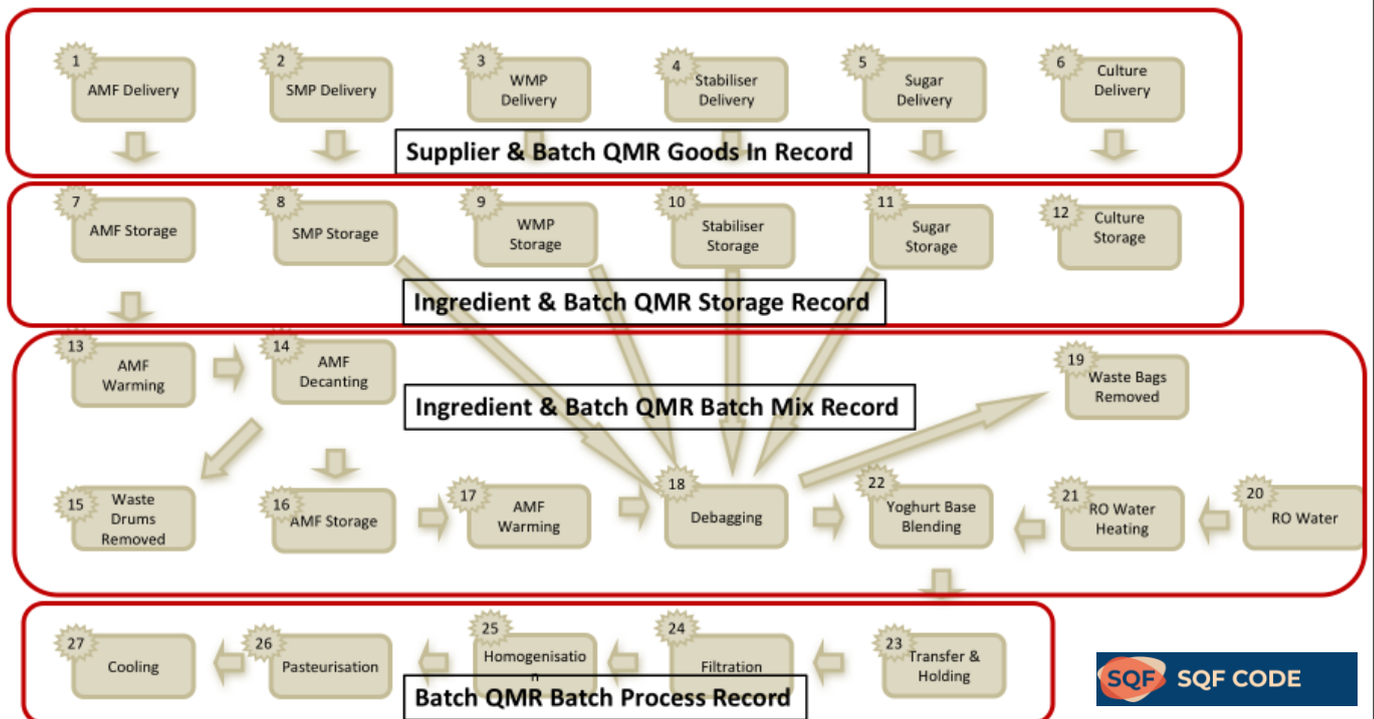
The effectiveness of the product trace system is reviewed at least annually as part of the product recall and withdrawal review. These exercises and any corrective actions are documented. Product trace tests are carried out on products from different shifts and for materials (including bulk materials) that are used across a range of products and/or products that are shipped to a wide range of customers.

Document Reference FS 2.6.2 Product Trace
Revision 0 11th March 2026
Owned by: Quality Manager
Authorized by: General Manager

Page 1 of 4 889 words English (United States) Text Predictions: On Accessibility: Unavailable Focus 65%

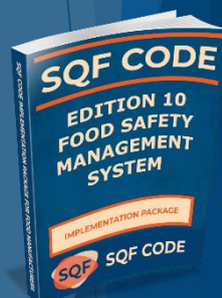
PowerPoint Slide Show - [FS 2.6.2A Traceability System Diagram]

FS 2.6.2A Traceability System Diagram

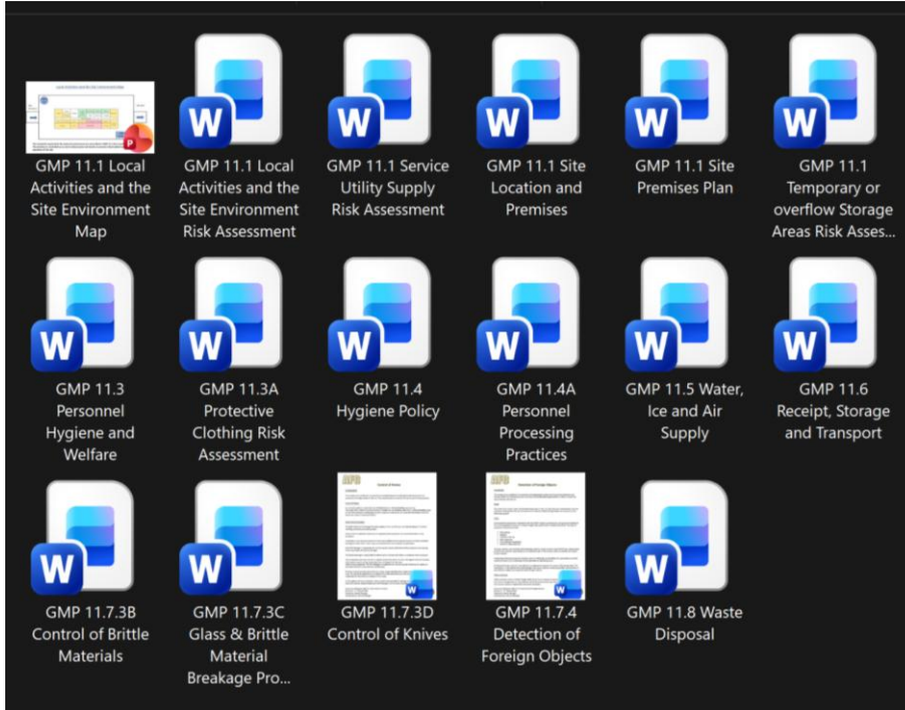
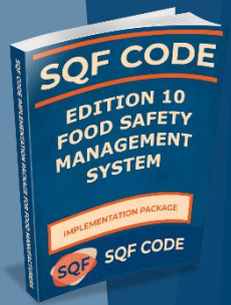


These are the procedure templates for Module 11: Good Manufacturing Practices for Processing of Food Products

- GMP 11.1.1 Site Location and Premises including:
 - Building Materials
 - Lighting and Light Fittings
 - Inspection/Quality Control Area
 - Dust, Insect, and Pest Proofing
 - Ventilation
 - Equipment and Utensils
 - Grounds and Roadways
- GMP 11.1A Site Premises Plan
- GMP 11.2.1 Repairs and Maintenance
- GMP 11.2.2 Maintenance Staff and Contractors
- GMP 11.2.3 Calibration
- GMP 11.2.4 Pest Prevention
- GMP 11.2.5 Cleaning and Sanitation
- GMP 11.3 Personnel Hygiene and Welfare including:
 - Hand Washing
 - Clothing and Personal Effects
 - Visitors
 - Staff Amenities
- GMP 11.3A Protective Clothing Risk Assessment
- GMP 11.4 Hygiene Policy
- GMP 11.4 Personnel Processing Practices
- GMP 11.5 Water, Ice and Air Supply
- GMP 11.6 Receipt, Storage and Transport including:
 - Receipt, Storage and Handling of Goods
 - Cold Storage, Freezing and Chilling of Foods
 - Storage of Dry Ingredients, Packaging, and Shelf Stable Packaged Goods
 - Storage of Hazardous Chemicals and Toxic Substances
 - Loading, Transport and Unloading Practices
- GMP 11.7.1 Separation of Functions & High-Risk Processes
 - GMP 11.7.1A Personnel High Risk Hygiene Barrier
 - GMP 11.7.2 Thawing of Food
- GMP 11.7.3 Control of Foreign Matter Contamination
 - GMP 11.7.3A Glass Policy
 - GMP 11.7.3B Control of Brittle Materials
 - GMP 11.7.3C Glass & Brittle Material Breakage Procedure
 - GMP 11.7.3D Control of Knives
- GMP 11.7.4 Detection of Foreign Objects
- GMP 11.8 Waste Disposal



There are also Risk Assessments as required in Module 11 of the new SQF Food Safety Code Edition 10



AutoSave ⏏ ↶ ↷ ⌵ GMP 11.1 Local Activities and the Site E... AC ⊞ ⌵

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Local Activities and the Site Environment Risk Assessment

Area Description	Description of Hazard	Likelihood	Severity	Significance	Details and Control Measures
Environment - Hardcore area	Pest Activity	1	2	2	Vegetation is minimized and controlled in this area
Environment - Vegetation area	Pest Activity	2	2	4	Vegetation is not controlled in this area. Extensive pest prevention measures and monitoring are in place on this side of the site.
Local River	Flooding	1	3	3	River has previously flooded to 2 m in the past 100 years. Site is on elevated hardcore 10m above the river level
Derelict Building - Hardcore area	Pest Activity	2	2	4	Pest prevention measures and monitoring are in place on this side of the site. Hardcore prevents easy access across to site
Sewage Works - 4km away	Flooding/Overflow	1	3	3	River has previously flooded to 2 m in the past 100 years. Sewage Works is on elevated hardcore 5m above the river level and has never flooded
Local Waste Dump - 5km away	Pest Activity	3	1	3	Pest prevention measures and monitoring are in place at Dump perimeter. No issue ever sourced from dump noted. Entrances are pest proofed and fly screening is in place on site.
Local Pig Farm - 6km	Surface flow of contaminated water	3	1	3	River has previously flooded to 2 m in the past 100 years. The pig farm can flood by run off is away from site.
Site Perimeter	Entry of local animals	1	2	2	Secure fencing and a gated entry is in place.

Revision 0 11th March 2026
 Owned by: Quality Manager
 Authorized by: General Manager

Page 2 of 2 332 words English (United States) Text Predictions: On Focus 65%

PACKAGE DOCUMENT EXAMPLES

AFC Site Location and Premises

6 Food facilities are located away from areas where wastes, either solid or liquid, cannot be removed effectively.

7 Measures established to maintain a suitable external environment are monitored and periodically reviewed. Periodic assessment of potential food safety impact from on and to local environment is performed.

8 Surroundings are kept neat and tidy and not present a hazard to the hygienic and sanitary operation of the premises. Paths from amenities leading to site entrances are required to be effectively sealed.

9 Site areas including footpaths, roads, yards, loading/unloading and parking areas are maintained so as not to present a hazard and have adequate drainage to prevent the accumulation of water.

10 The grounds, area surrounding the premises storage facilities, machinery, and equipment are maintained to minimize dust and kept free of waste, accumulated debris or standing water so as not to attract pests and vermin or present a food safety hazard to the sanitary operation of the site.

Ceilings

1 All ceilings are solid and not hollow

2 All ceilings are fire resistant

3 All ceilings and their finishes are impervious and non-absorbent, washable and easily cleaned, non-contaminating and non-staining

4 Ceilings and overhead fixtures are constructed to minimize the buildup of dirt and constructed and maintained to prevent the contamination of products

5 Drop ceilings have adequate access to the void for cleaning and pest management and are checked and cleaned regularly

6 Where drop ceilings are not used, cleaning regimes and inspections are in place to check for dust on ledges, loose fittings, glass windows, light fittings, or other areas where dust can accumulate and fall onto product

Floors

1 Floors are made of specified materials which are durable, impact resistant, impervious and non-absorbent, washable and easily cleaned, non-contaminating, non-tainting and have an even and regular surface

2 Floors are constructed with gradients to allow adequate drainage and cleaning. Where floor drainage is not available, plumbed options to handle and remove overflow or wastewater are put in place

3 Drains are constructed and located so they can be easily cleaned and do not present a hazard

4 Waste trap systems are located away from any food handling area and entrants to the premises

Internal Walls

1 All internal walls are solid and not hollow

2 Internal walls are damp proofed and fire resistant

Document Reference GMP 11.1 Site Location and Premises
Revision 0 13th March 2026
Owned by: Quality Manager
Authorized by: General Manager

AFC Cleaning and Sanitation

Correct dilution and temperature of Chemicals
Methods used to confirm the correct concentrations of detergents and sanitizers
Contact time for Chemicals
Method of Cleaning
Any precautionary measures
Frequency of cleaning
Personal responsible for cleaning

The company operates a clean as you philosophy which is briefed to all staff and monitored by department managers to ensure all personnel keep their areas in a clean and tidy state. Cleaning tools and equipment are of hygienic design and maintained in a condition which does not represent a risk to the product.

Suitably equipped areas are designated for cleaning product containers, knives, cutting boards and other utensils and for cleaning of protective clothing used by staff. These cleaning operations are controlled so as not to interfere with manufacturing operations, equipment or product. Racks and containers for storing cleaned utensils are provided.

A register is maintained of all Cleaning Chemicals approved for use on site. A chemical control sheet is in place for each chemical used on site which includes details the management of use, handling and storage of non-food chemicals including:

- Approved supplier
- Chemical data and safety sheets
- Suitability for use as appropriate to use
- Instructions for the avoidance of use of chemicals with strong aromas in manufacturing and storage areas
- Identification of chemicals
- Segregated and secure storage areas
- Use by trained personnel

Cleaning chemicals are fit for purpose, suitably labelled according to regulatory requirements, stored in closed containers and used in accordance with manufacturers' instructions. Detergents and sanitizers must be suitable for use in a food manufacturing environment. All cleaning chemicals and equipment are clearly identified and segregated. Detergents and sanitizers are controlled in accordance with manufacturers' instructions, stored in containers that are suitable for use, and clearly identified. Mix concentrations are verified, records maintained and chemical usage monitored.

Chemicals are applied and stored according to label directions. Empty chemical containers are disposed of according to label directions and regulatory requirements; unused obsolete chemicals are secured until collected by the supplier and disposed of as per regulatory requirements.

Document Reference GMP 11.2.5 Cleaning and Sanitation
Revision 0 13th March 2026
Owned by: Quality Manager
Authorized by: General Manager

AFC Pest Prevention

The contracted service provides:

- Monthly site visits and inspections including service records describing current levels of pest activity and recommendations for taking Corrective Actions.
- Inspections including the periphery and internal and external buildings
- The provision of a plan/diagram of the site showing the identification, location, number and type of all pest control monitoring and prevention measures
- Flying insect controls including fly killing units
- Emergency 24 hour call-out service
- Quarterly biologist inspection reports, visit and trend reports with recommendations
- A record of pest sightings and a trend analysis of the frequency of pest activity to target pesticide applications
- Current copy of the certification of insurance that specifies the liability coverage
- Disposal of unused pest control chemicals and empty containers in accordance with regulatory requirements
- Spill control materials and procedures
- Safety Data Sheet information to ensure proper usage of pesticide chemicals.

Both the contract and service agreement information are held in the Pest Control File which is managed by the Quality Manager who has overall responsibility for pest control on site.

Before agreeing to a contract, the Quality Manager verifies that the pest control contractor is qualified. Copies of training records and qualifications are held in the pest control file for each person who performs pest management services on site. At the start of the contract a detailed survey of the entire facility is completed by a qualified Field Biologist and the results are documented and used to determine placement of monitoring devices.

Exterior Bait Stations

Exterior rodent bait stations are set up to deter rodents from entering the facility. Based on the detailed facility survey, exterior bait stations are placed along the foundation walls on the exterior of the facility and along the site boundaries. Exterior bait stations containing rodenticides are tamper resistant, anchored in place, locked, and labelled. All exterior bait stations are inspected at least monthly. The bait stations are checked more often when activity levels increase. Bait stations are secured inside bait stations, in good condition, and replaced as needed. Bait stations are placed at intervals of 15 m although areas of high rodent activity may have a higher concentration of bait stations.

Interior Monitoring

Based on the detailed Field Biologist survey, interior monitoring devices are placed in strategic sensitive areas specific to the rodent species, and other areas of pest activity, including:

- Raw material warehouse

Document Reference FS 11.2.4 Pest Prevention
Revision 0 13th March 2026
Owned by: Quality Manager
Authorized by: General Manager

Supplementary HACCP Documents, Guidance and Tools

The HACCP Calculator and Instructions

HACCP Calculator based SQF Edition 10 requirements and the latest version of CODEX General Principles of Food Hygiene Chapter Two HACCP System and Guidelines for its Application including a new Decision Tree.

AutoSave | SQF Edition 10 HACCP Calculator | Saved to this PC

File Home Insert Draw Page Layout Formulas Data Review View Automate Help

HACCP CALCULATOR SQF Edition 10 Version

Step Number	Step Name	Hazard Category	Hazards Identified	Specific Details about the Hazard	Existing GMPs which assist in controlling the Hazard	Control Measure	P	r	r	S	S	S	Q	Q	Q	Q	C	G	M	P	Critical Limits	Monitoring Procedures	Corrections & Corrective Action
							Pre	ro	o	Se	Si	Q	Q	Q	Q	C	G	M	P				
1	AMF Delivery	Biological	Bacteria (spore-forming) General		No GMPs assist in Control of Hazard	Storage 1 - 5 °C							N	Y	N	N					No Contamination Always under cover	Supervision by Warehouse Manager	Retrain Staff. Inspect delivery for contamination. Reject if contaminated
11	1	AMF Delivery	Biological	Bacillus cereus	No GMPs assist in Control of Hazard	Storage 1 - 5 °C	3	3	3				N	Y	N	N							
12	1	AMF Delivery	Biological	Bacteria (spore-forming) General	No GMPs assist in Control of Hazard	Storage 1 - 5 °C	3	3	3				N	Y	N	N							
14	1	AMF Delivery	Biological	Bacteria (spore-forming) General	No GMPs assist in Control of Hazard	Storage 1 - 5 °C	3	3	3				N	Y	N	N							
15	1	AMF Delivery	Biological	Bacteria (spore-forming) General	No GMPs assist in Control of Hazard	Storage 1 - 5 °C	3	3	3				N	Y	N	N							
16	1	AMF Delivery	Chemical	Antibiotics	No GMPs assist in Control of Hazard	Storage 1 - 5 °C	2	2	4														
17	1	AMF Delivery	Allergens	Soya	No GMPs assist in Control of Hazard	Storage 1 - 5 °C	1	1	1														
18	1	AMF Delivery	Physical	Glass	No GMPs assist in Control of Hazard	Storage 1 - 5 °C	3	2	6														
19	1	AMF Delivery	Radiological	Cesium-134	No GMPs assist in Control of Hazard	Storage 1 - 5 °C	3	1	3														
20	1	AMF Delivery	Bacteria (spore-forming) General	11.3 Cleaning and Sanitation and Pest Prevention	11.3 Cleaning and Sanitation and Pest Prevention	Storage 1 - 5 °C	3	2	6														

Decision Tree **

- STOP Not a CCP
- N Go to next Question
- Y Go to next Question
- Y That next step is a CCP
- N Modify ****
- Y This is a CCP

* Consider the significance of the hazard (i.e., the likelihood of occurrence in the absence sufficiently controlled by prerequisite programs such as GHPs. GHPs could be routine GHP monitoring.

** If a CCP is not identified at questions 2-4, the process or product should be modified

*** Consider whether the control measure at this step works in combination with a control should be considered

**** Modify the step, process or product

Process Flow | HACCP Calculator | HACCP Plan | HACCP Validation | Good Manufacturing Practices | Control Mea

Ready | Accessibility: Investigate | 75%

The HACCP Calculator is a tool that allows you to present your Hazard Analysis in a clear and logical manner.

From the process flow you select hazards and assess them to identify significant hazards.

You then use the in-built CODEX Decision Tree questions to determine your Critical Control Points.

Following that the HACCP Calculator assists in developing Food Safety Plans to control Significant Hazards.

Decision Tree

- STOP Not a CCP
- Go to next Question
- That next step is a CCP
- Modify ****
- This is a CCP

Hazard List

Step Number	Step Name	Hazard Category	Hazards Identified	Specific Details about the Hazard	Existing GMPs which assist in controlling the Hazard	Control Measure	C	L	M	P	C	C	A	R	H	V
1	AMF Delivery	Biological	Bacteria (spore-forming) General		No GMPs assist in Control of Hazard	Storage 1 - 5 °C	3	3	3	3	3	3	3	3	3	3
1	AMF Delivery	Biological	Bacillus cereus		No GMPs assist in Control of Hazard	Storage 1 - 5 °C	3	3	3	3	3	3	3	3	3	3
1	AMF Delivery	Biological	Bacteria (spore-forming) General		No GMPs assist in Control of Hazard	Storage 1 - 5 °C	3	3	3	3	3	3	3	3	3	3
1	AMF Delivery	Biological	Bacteria (spore-forming) General		No GMPs assist in Control of Hazard	Storage 1 - 5 °C	3	3	3	3	3	3	3	3	3	3
1	AMF Delivery	Biological	Bacteria (spore-forming) General		No GMPs assist in Control of Hazard	Storage 1 - 5 °C	3	3	3	3	3	3	3	3	3	3
1	AMF Delivery	Chemical	Antibiotics		No GMPs assist in Control of Hazard	Storage 1 - 5 °C	2	2	2	2	2	2	2	2	2	2
1	AMF Delivery	Allergens	Soya		No GMPs assist in Control of Hazard	Storage 1 - 5 °C	1	1	1	1	1	1	1	1	1	1
1	AMF Delivery	Physical	Glass		No GMPs assist in Control of Hazard	Storage 1 - 5 °C	3	2	2	2	2	2	2	2	2	2
1	AMF Delivery	Radiological	Cesium 134		No GMPs assist in Control of Hazard	Storage 1 - 5 °C	3	1	1	1	1	1	1	1	1	1
2	SMP Delivery	Bacteria (spore-forming) General		11.3 Cleaning and Sanitation and Pest Prevention	No GMPs assist in Control of Hazard	Storage 1 - 5 °C	3	2	2	2	2	2	2	2	2	2
2	SMP Delivery	Bacteria (spore-forming) General		11.3 Cleaning and Sanitation and Pest Prevention	No GMPs assist in Control of Hazard	Storage 1 - 5 °C	2	3	3	3	3	3	3	3	3	3
2	SMP Delivery	Bacteria (spore-forming) General		11.3 Equipment and Maintenance	No GMPs assist in Control of Hazard	Storage 1 - 5 °C	2	3	3	3	3	3	3	3	3	3
2	SMP Delivery	Bacteria (spore-forming) General			No GMPs assist in Control of Hazard	Storage 1 - 5 °C	2	3	3	3	3	3	3	3	3	3
2	GMP Delivery	Bacteria (spore-forming) General			No GMPs assist in Control of Hazard	Storage 1 - 5 °C	1	3	3	3	3	3	3	3	3	3

HACCP Calculator Instructions

1 HACCP Calculator Instructions SQF 10 & CODEX

2 HACCP Calculator Instructions

3 SQF HACCP Application

A food safety plan shall be prepared in accordance with the twelve steps identified in the latest version of Codex Alimentarius Commission's General Principles of Food Hygiene.

CODEX ALIMENTARIUS
INTERNATIONAL FOOD STANDARDS ORGANIZATION

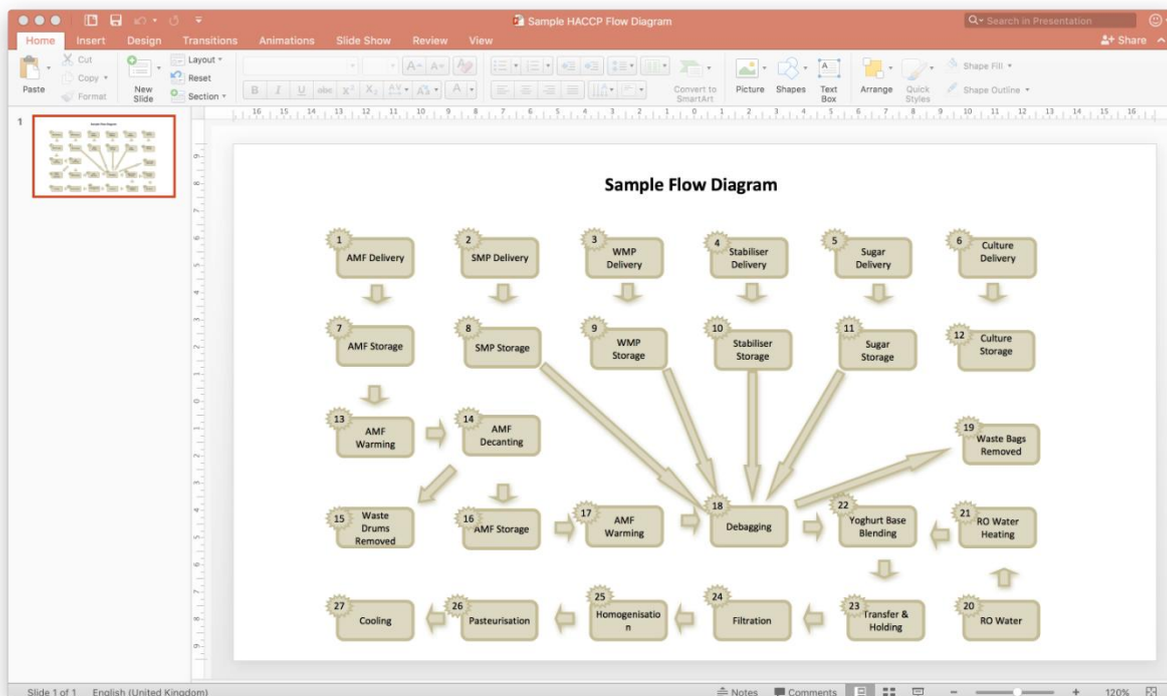
4 CODEX HACCP Application

The HACCP system and guidelines for its application are defined by the Codex Alimentarius Commission in the CODEX Recommended International Code of Practice General Principles of Food Hygiene CXG 1-1969. Adopted in 1969. Amended in 1999. Revised in 1997, 2003, 2020, 2022. Editorial corrections in 2011.

CODEX ALIMENTARIUS
INTERNATIONAL FOOD STANDARDS ORGANIZATION

Supplementary HACCP Documents, Guidance and Tools

Useful additional HACCP Documents are included



There are supplementary HACCP document templates including Flow Diagrams, Product Description, a Hazard Analysis Prompt, an example Critical Control Procedure and various HACCP Records.

AFC Pasteurizer Log Sheet

DATE: _____

Product:	Tank:	Product:	Fat %:	Total Solids:	Temp. (°C):	QC. Sign:
Feed Tank:	Fill Tank:					
Volume:						
Production Start Time:	Production End Time:	CIP Start/End Time:				

PARAMETERS	LIMITS	UNITS
Flow Rate (CCP Maximum 5250)	5000-5250	L/h
Pre-heater In Temperature	45 - 55	°C
Pasteurization Temp. (Homo in Temp.)	82 ± 2	°C
Pasteurizer Out Press.	2.8-3.0	PI
Homo in Press.	1.8-2.0	PI
Pressure Difference (CCP)	Minimum 0.8	PI
End Holding Temp. (CCP)	Min. 77.0	°C
Product Outlet Temp. (CCP)	< 5	°C
Homo Press. (1st/ 2nd Stage)	175/ 50	Bar
Homo Pressure (Total)	225	Bar
Glass & Perspex Items Check & Sign	Intact/No Cracks	
Sterilization Temperature	82 ± 2	°C
Diversion Test Before Production	Minimum 77	°C
Record Diversion Temperature & Sign		

Operator Name & Sign: _____ Supervisor Sign: _____

Document Reference Pasteurizer Log Sheet PAS 001
Revision 0 1st August 2022
Owned by: Production Supervisor
Authorized by: Production Manager

AFC Ice Cream Pasteurization Procedure

PARAMETERS	LIMITS	UNITS
Preheater in Temp.	45 - 50	°C
Holding time (CCP) Min. 15 seconds	Min 15	s
Pasteurizer in Press.	0.5 - 1.0	Bar
Pasteurization Temp.	73 ± 1	°C
End Holding Temp. (CCP) Min. 72.0 °C	73 ± 1	°C
F. Cooler Out Flow Rate	5.0-5.25	m ³ /h
Milk Outlet Temp.	4 ± 2	°C
Product Outlet Overpressure	> 1.0	Bar
Homo Press. (1st/ 2nd Stage)	150/50	Bar

Ensure that the Pasteurization Temperature is 73 ± 1 °C (Min. 72 °C) and the holding time is a minimum of 15 seconds.

During processing, to change to another Ice Cream Tank put the pasteurizer on recirculation, change to the required tank then press forward flow.

When the product finishes flush the pasteurizer with water. Record the Volume Processed, Processing Time & Production End Time.

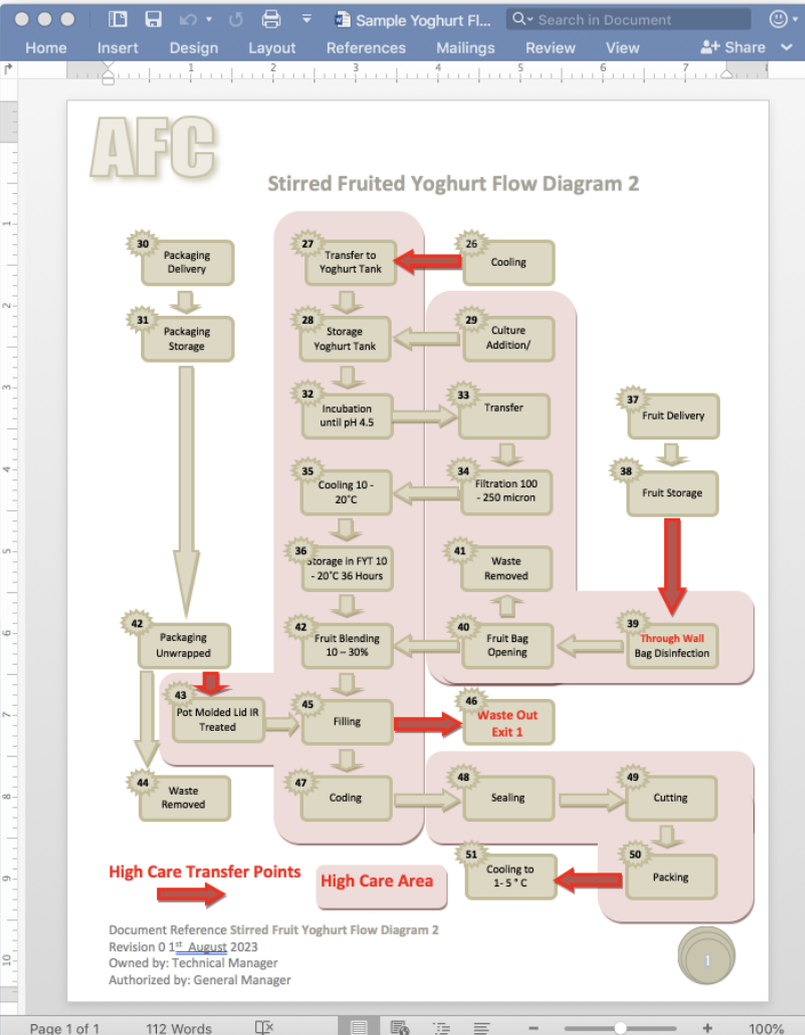
After rinsing proceed to Clean in Place. Record the CIP Start & End Times.

IF ANY PROCESS PARAMETERS ARE OUT OF SPECIFICATION DO NOT CONTINUE TO PROCESS, PUT THE PASTEURISER ON RECIRCULATION AND CONTACT THE PASTEURISER SUPERVIZER IMMEDIATELY.

REFERENCES

1kg Ice Cream Specification SPEC 1
FSR 1 Pasteurizer Log Sheet

Document Reference Ice Cream Pasteurization Procedure FS 1
Revision 0 1st August 2022
Owned by: Pasteurizer Supervisor
Authorized by: Production Manager

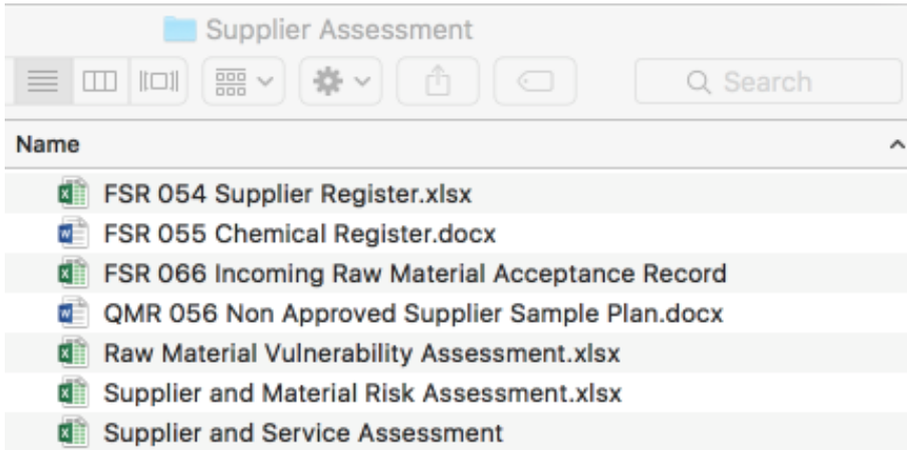


AFC Hazard Analysis Prompt

Hazard Analysis Prompt	Answers in Detail
Are the raw materials, ingredients or food contact packaging likely to have chemical, biological or physical hazards present?	
Are there any characteristics in the composition of the food during which can prevent a hazard? E.g. Preservatives, pH, Water Activity	
Does the food permit survival or multiplication of pathogens and at which stages?	
Does the process include a controllable step that destroys pathogens or their toxins? (Consider spores) Is it possible the product could be subject to recontamination?	
Is product contamination (consider direct and indirect contamination) with hazardous microbiological organisms from equipment, process environment or personnel likely to occur?	
Is product contamination (consider direct and indirect contamination) with hazardous chemical substances from equipment, process environment or personnel likely to occur?	
Is product contamination (consider direct and indirect contamination) with hazardous physical objects from equipment, process environment or personnel likely to occur?	
Is it likely that the food contains viable spore forming pathogens?	
Is it likely that the food contains viable non-spore forming pathogens?	
What is the normal microbial content of the food stored under proper conditions?	
Does the microbial population increase during the time the food is stored before consumption?	
Does that increase in microbial population alter the safety of the food?	
Does the layout of the facility provide an adequate separation of raw materials from ready-to-eat foods?	
Will the equipment provide the time and temperature control that is necessary to meet critical limits?	
Is the equipment reliable or is it prone to frequent breakdowns?	

Document Reference HM 8 Hazard Analysis Prompt
Revision 0 11th March 2026
Owned by: Quality Manager
Authorized by: General Manager

Supplementary Supplier Assessment Documents and Tools



There are assessment tools and sample records

Raw Material Vulnerability Assessment

Raw Material Vulnerability Assessment Calculator

A poor harvest may restrict availability and may increase the potential for adulteration, Sophistication of routine testing to identify adulterants (if testing within the supply chain is comprehensive and focused on potential fraud issues, then the likelihood is less), Country of origin, length and complexity of the supply chain

Score	Material Category Rating
5	No raw material with recent reports of adulteration published by regulatory authorities - action or monitoring is required to ensure only genuine material is purchased.
4	The material that provides an attractive target for potential adulteration - some action and/or monitoring is required to ensure only genuine material is purchased.
3	Medium - a material that may be adulterated - action is required to ensure only genuine materials are purchased.
2	This material is unlikely to be a target for substitution or adulteration, however a re-assessment may be necessary if new information becomes available.
1	Negligible - no further action required as the material is extremely unlikely to be a target for food fraud.

Supplier Number	Supplier	Materials Supplied	Material Category	Historical evidence of substitution or adulteration	Economic factors which may make adulteration or substitution more likely	Ease of access to raw materials through the supply chain	Sophistication of routine testing to identify adulterants	Stature of the Raw Material	Current Controls in Place	Primary Control	Secondary Control
1	A	Chocolate Topping	Final Ingredient						Supplier Audit every 6 months	Certificates of analysis from raw material suppliers	Positive Release by Site prior to Use
2	B	Flour for Baking	Raw Ingredient						Supplier Audit every 2 Years	Raw material testing	Supply chain audits
3	C	Contract Scones	Contract Packager						Certification to GFSA Approved Standard	Mass balance exercises at the raw material supplier	Stamp evidence on Incoming raw materials
4	D	Cake Tray	Contract Packaging						Supplier Assurance Questionnaire	Raw material testing	CDC with each Delivery
5	E	Cardboard Box	Non-Contact Packaging						Supply to Contract Specification	Certificates of analysis from raw material suppliers	
6	F	Non-Contact Packaging							Supplier Audit every 6 months	Supplier Audit every 6 months	

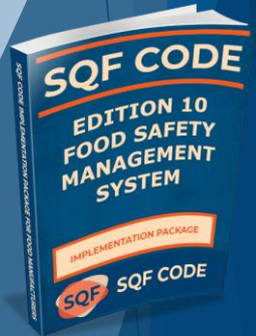
Supplier and Material Risk Assessment

Allergens and possible allergen contamination

Supplier & Material Risk Calculator

Score	Supplier Category Rating	Severity of Risk	Risk Score	Rating	What should I do?
5	Final Ingredient/Contract Packager	Catastrophic - death or large number of serious injuries	25	Extreme	Clear Surveillance of Supplier and Material Required
4	Raw Ingredient/High Risk Service	Major - serious injury, extensive injuries	16 - 20	High	Supplier and Material/Service Monitoring Required
3	Contract Packaging	Moderate - medical treatment required	9 - 15	Moderate	Material/Service Monitoring Required
2	Non-Contact Packaging	Minor - first aid treatment required	< 9	Low	Prerequisites on Goods in/Service Provision Sufficient
1	Low Risk Service	Minor - no injuries			

Supplier Number	Supplier	Materials/Service Supplied	Supplier Category	Identify the Risks	Details of Hazard or Risk	Use the Current Controls in Place	Supplier Control	Material Control	Supplier Control
1	A	Chocolate Topping	Final Ingredient	Microbiological contamination	Not Further Processed on Site	5	Supplier Audit every 6 months	Positive Release by Site prior to Use	
2	B	Flour for Baking	Raw Ingredient	Chemical contamination	Further Processed on Site	4	Supplier Audit every 2 Years	Certification to GFSA Approved Standard	
3	C	Contract Scones	Contract Packager	Chemical contamination	None Currently	5	Supplier Audit every 6 months	Certification to GFSA Approved Standard	
4	D	Cake Tray	Contract Packaging	Ins and possible allergen contamination	Packaging Reused and Inverted	3	Supplier Assurance Questionnaire	Supplier Assurance Questionnaire	
5	E	Cardboard Box	Non-Contact Packaging	Possible substitution of fraud	No access to Production Facility	1	Supplier Assurance Questionnaire	CDC with each Delivery	



Supplement to Product Inspection, Onsite Product Testing and Laboratory Analysis

In addition to FS 2.4.4 Product Inspection, Testing and Analysis Procedure, a comprehensive Laboratory Quality Manual compliant with the requirements of ISO 17025 is provided in Microsoft Word format.

The Laboratory Quality Manual includes template records, procedures and product sampling plans.

AFC Laboratory Quality Manual

CONTENTS

1. Introduction
2. Quality System
3. Organization and Management
4. Personnel
5. Laboratory Accommodation and Environment
6. Personnel Hygiene
7. Confirmation of Work and Client Requirements
8. Handling Test Items
9. Test Methods
10. Bench Practices
11. Assuring Quality of Results
12. Equipment, Calibration and Measurement Traceability
13. Calibration Standards / Reference Materials
14. Reporting Test Results
15. Records
16. Purchase of Outside Services, Supplies and Laboratory Consumables

Document Reference FS 2.4.4A Laboratory Quality Manual
Revision 0 1st March 2026
Owned by: Laboratory Supervisor
Authorized by: Quality Manager

AFC Factory Sample Plan

Sample	Point	Test / Inspection	Frequency	Standard	Method Ref	Spec Ref	Record / Log Ref
Liquid Ingredient 1	Tank	% AW	Each Load F & R	Max 85%	AP 001	LSP 001	LBR 001
		% Fat		> 5%	AP 002	LSP 001	LBR 001
		% Acidity		0.1 - 0.2	AP 003	LSP 001	LBR 001
		Enterobacteriaceae		< 10/ml	MP 001	LSP 001	LBR 001
		TVC		< 10,000cfu/g	MP 002	LSP 001	LBR 001
		Phosphatase		Pass	AP 004	LSP 001	LBR 001
		Smell		Fresh Normal	AP 005	LSP 001	LBR 001
		Taste		Fresh Normal	AP 006	LSP 001	LBR 001
Ingredient in Storage	Silo	Age	Each Load F & R	< 48 Hours	AP 003	LSP 001	LBR 001
		% Acidity		0.1 - 0.2	AP 003	LSP 001	LBR 001
		Smell		Fresh Normal	AP 005	LSP 001	LBR 001
		Taste		Fresh Normal	AP 006	LSP 001	LBR 001
Ingredient 3	Tank	% Fat	Each Flow Box	10% +/- 1%	AP 002	LSP 001	LBR 001
		% Acidity		0.10 - 0.20	AP 003	LSP 001	LBR 001
		Temperature		< 7 °C	AP 007	LSP 001	LBR 001
		Enterobacteriaceae		< 10/g	MP 001	LSP 001	LBR 001
		Phosphatase		Pass	AP 004	LSP 001	LBR 001
		Smell		Fresh Normal	AP 005	LSP 001	LBR 001

Document Reference Factory Sample Plan LAB 007
Revision 0 1st August 2023
Owned by: Laboratory Supervisor
Authorized by: Quality Manager

Supplementary Laboratory Quality Manual

Search

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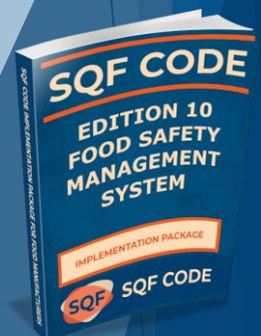
- LABR 001 Laboratory Audit Form.docx
- LABR 002 Laboratory Training Form.docx
- LABR 003 Laboratory Autoclave Record.docx
- LABR 004 Microbiological Sample Plan.docx
- LABR 005 Filler Sample Plan.docx
- LABR 006 QA Sample Plan.docx
- LABR 007 Factory Sample Plan.docx
- LABR 007 Factory Sample Plan.xlsx
- LABR 008 Daily Balance Calibration Sheet.docx
- LABR 009 Laboratory Exception Report.docx
- LABR 010 QC Online Check Sheet.docx
- LPOL 001 Laboratory Quality Policy.docx
- LPPRO 001 Laboratory Operating Procedure for the Autoclave.docx
- MICRO 001 Enumeration of Total Viable Counts.docx

AFC Laboratory Daily Exception Report

Date:

Area	RO Water	Process Checks	Fresh		Packing		
			Filler 1	Filler 2	1	2	3
Enteros							
ATP Swab/Rinse							
TVC							
AKQ							
Shelf Life							
Chemical Analysis							
Weight/Volume							
CIP Checks	Caustic Strengths Target 1.8 – 2.2%	Acid Strengths Target 1.3 – 1.7%	Report any issues with each CIP set				
CIP 1							
CIP 2							
CIP 3							
CIP 4							

Document Reference Laboratory Daily Exception Report
Revision 0 1st August 2023
Owned by: Laboratory Manager
Authorized by: Quality Manager



Supplementary Allergen Management Tools

FS 2.8.1 Allergens Management is a comprehensive Allergen Management Procedure which is supplemented by Allergen Management Tools and other useful Allergen Control Documents

The screenshot displays three pages of a Microsoft Word document titled "FS 2.8.1 Allergen Management".

- Page 1:** Introduction. The company recognizes the serious repercussions of allergic reactions and therefore takes every precaution to prevent this happening. The company has established an Allergen Control System (ACS) which is maintained as part of the operational programmes in order to meet the requirements of the Food Safety Management System and ensure the safe production of products. It lists various allergen categories like Peanuts, Tree Nuts, Cereals containing Gluten, Milk, Eggs, Fish, Crustacean Shellfish, Soy, Sesame seeds, Celery/celeric, Mustard, Lupin, Sulphur dioxide and sulphites.
- Page 2:** Allergen Control System. An Allergen Control System has been implemented to control allergens on site and to minimize the risk of the unintentional inclusion of allergens in products. It details the steps in the Allergen Control System, including identification of raw materials, ingredients, and processing aids, and the identification of products containing allergens.
- Page 3:** Identification of Relevant Allergens as per Legislation and Customer Requirements. Relevant allergens and acceptable levels are prescribed by legislation, customer requirements and industry code of practice. It includes a table for "Allergen Management Tool - Allergen List" and a table for "Identification of Ingredients with Allergen Content/Possible Allergen Content".

- FS 2.8.1 Allergen Management.docx
- FS 2.8.1A Allergen Management Tool.xlsx
- FS 2.8.1B Allergen Clean Validation.docx
- FS 2.8.1C Allergen Clean Verification.docx
- FS 2.8.1D Ingredient Allergen - Color Coding EU.docx
- FS 2.8.1D Ingredient Allergen - Color Coding USA
- FS 2.8.1E Allergens.docx
- FS 2.8.1F Allergen Management Records

- Finished Product Allergen Summary.docx
- Supplier Ingredient Allergen Analysis Form.docx
- Allergen Warning Label Color Coding Summary.docx
- Raw Material Allergen Summary Form.docx
- Allergen Warning Label - Sesame seeds.docx
- Allergen Warning Label - Soybeans.docx
- Allergen Warning Label - Wheat.docx
- Allergen Warning Label - Peanuts.docx
- Allergen Warning Label - Tree Nuts.docx
- Allergen Warning Label - Crustacean Shellfish.docx
- Allergen Warning Label - Fish.docx
- Allergen Warning Label - Eggs.docx
- Allergen Warning Label - Milk.docx

The screenshot shows a Microsoft Word document titled "FS 2.8.1D Ingredient Allergen Management - Color Coding USA". It includes a legend for allergen color coding:

- Milk (Yellow)
- Eggs (Green)
- Fish (Blue)
- Crustacean Shellfish (Grey)
- Tree Nuts (Orange)
- Peanuts (Red)
- Wheat (Purple)
- Soybeans (Black)
- Sesame seeds (Brown)

The document footer indicates: Document Reference FS 2.8.1D Appendix Ingredient Allergen Management - Color Coding USA, Revision 0 1st August 2023, Owned by: Quality Manager, Authorized by: General Manager.

Product Development Tools

FS 2.3.1 Product Design & Development Procedure is supported by supplementary Product Design & Development documents and forms and a Product Development Plan

The screenshot shows a Microsoft Word document titled "NPD 1 Product Development Plan" with the "AFC" logo. The document contains a "Product Development Plan" table with the following structure:

Stage	Responsibility	Date	Signed
STAGE 3: Approval of Kitchen Product			
- Product Approval by Customer			
- Reference sample saved			
- Full raw material Specification & Supplier Questionnaire or audit, checked, completed and to be signed by both parties			
- Project Schedule updated			
- Handover to process development			
STAGE Complete & Authority to Move to Next Stage	<u>Yes/No</u>	<u>Date</u>	<u>Signed</u>

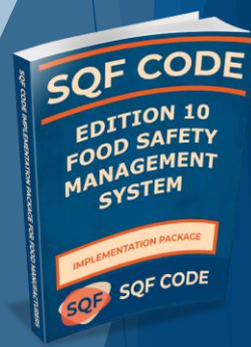
Below the table, the document reference information is provided:

Document Reference NPD 1 Product Development Plan
 Revision 0 11th March 2026
 Owned by: Development Manager
 Authorized by: Quality Manager

The document footer shows "Page 2 of 9", "661 words", "English (United States)", "Text Predictions: On", "Accessibility: Unavailable", "Focus", and "100%" zoom.

The screenshot shows a file explorer window titled "Supplementary Product Development Tools" with a search bar and a list of files:

- FPSPEC 001 Whole Milk Summer F...io Yoghurt 100g Specification.docx
- FPSPEC 002 Whole 3.5% UHT Milk Specification.docx
- FPSPEC 003 1.5% Natural Set Yoghurt Specification.docx
- NPD 001 Product Development Plan.docx
- NPD 002 Product Development Brief Sign Off Form.docx
- NPD 003 Artwork Approval Form.docx
- NPD 004 Market Review Form.docx
- NPD 005 Project Request Form.docx
- NPD 006 Development Recipe Sheet.docx
- NPD 006 NPD Costing Form.docx
- NPD 007 Taste Panel Form.docx
- NPD 008 Factory Trial Assessment Form.docx
- RMS 001 Milk Powder Specification.docx
- RMS 002 Refined White Sugar Specification.docx
- RMS 003 Cocoa Powder Specification.docx
- RMS 004 Chocolate Specification.docx
- RMSP 001 Fruit Conserve Sample Plan.docx



Product Development Tools

AutoSave FS 2.3.1 Product Development - Compatibility Mode

File Home Insert Draw Design Layout References Mailings Review View Help

Comments Editing Share

AFC Product Development

Following the HACCP system review the Quality Manager and the Operations Manager authorize the process changes and a completed Process Change Approval form is held on file.

The Quality Manager and Operations Manager are responsible for ensuring that the site is capable of accommodating the new process or product and maintaining acceptable levels of quality and food safety.

A new specification will be created, defining raw materials, packaging and suppliers, Food Safety Plan, GMPs, in-process and finished product specifications and artwork, labelling and coding details. The New Product Development Manager confirms that the packaging complies with specifications, relevant food safety legislation and is fit for purpose. For all product claims the New Product Development Manager validates the product formulation and product process are capable of meeting the product claim prior to launch. Finished product specifications are agreed and authorized by the Development Manager and Quality Manager. The Quality Manager verifies that the labelling is adequate in informing customers of any critical allergen ingredients, relevant nutritional contents, storage, preparation and serving instructions and that the customer information meets legislation for the destination country.

The Quality Manager and Product Development Manager are responsible for ensuring that products that have not been in production for more than a year, or products that were previously discontinued are regarded as new products and are required to follow the product development procedure for approval prior to relaunch.

References

Appendix 1 NPD Stages
Product Development Folder

Document Reference FS 2.3.1 Product Development
Revision 0 11th March 2026
Owned by: Product Development Manager
Authorized by: Quality Manager

AFC Product Development

Appendix 1 NPD Stages

STAGE 1: Product Brief

- Product Brief supplied to NPD
- Critical path generation

STAGE 2: Kitchen work stage

- Specification sent for New Ingredients
- Preliminary Specification Checked and signed off
- Raw Material evaluated by Quality against the Spec
- Food contact packaging confirmed as suitable for intended use by Supplier
- Initial Product costing done
- All recipes documented

STAGE 3: Approval of Kitchen Product

- Product Approval by Customer
- Reference sample saved
- Full raw material Specification & Supplier Questionnaire or audit, checked, completed and to be signed by both parties
- Project Schedule updated
- Handover to process development

STAGE 4: Factory trials

- Food Safety Team approve factory trial
- Sign off / approval of any new equipment

Document Reference FS 2.3.1 Product Development
Revision 0 11th March 2026
Owned by: Product Development Manager
Authorized by: Quality Manager

AFC Product Development

- Positively release ingredients for factory trial
- Trial appraisal form/ report completed
- Yield Analysis
- Statistical Analysis
- Set Points Established
- Process capability study completed
- Initial Standards Tolerances set
- Scale up to production level
- Initial Packaging Trials undertaken
- Transit trials to store
- Re-cost Product if required

STAGE 5: Approval of Factory Product & Product Analysis

- Consumer panel if required
- Reference sample saved
- Samples sent for Nutritional
- Nutritional Results received
- Food Safety (HACCP) drawn up & verified
- Verification/Validation of the cooking instructions
- Samples sent for micro Shelf-life from 3 factory trial runs
- Micro shelf-life results forwarded to Quality Manager
- Organoleptic shelf-life started from 3 factory trial runs

Document Reference FS 2.3.1 Product Development
Revision 0 11th March 2026
Owned by: Product Development Manager
Authorized by: Quality Manager

Page 4 of 8 1919 words English (United States) Text Predictions: On Accessibility: Unavailable Focus 65%

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Calibri (Body) 14

AFC Whole Milk Summer Fruit Bio Yoghurt 100g

Manufacturing Site	
Contact Details	
Telephone	
Fax	

Product Description	
A whole milk stirred fruited bio yogurt with a creamy mixed berry flavour	
Organoleptic	
Appearance	Mauve in colour, smooth, shiny yoghurt with blackberry & raspberry pieces
Aroma	A fresh fruity mixed berry aroma
Flavour	Sweet creamy fresh mixed berry flavour with a slight lactic note

Ingredients	
Potable Water, Whole Milk Powder, Sugar, Blackberries (3.75%), Raspberries (3.75%) Summer Fruit Syrup [(water, glucose syrup, thickeners (modified starch, carrageenan), black carrot juice concentrate, woodberry flavor, sodium citrate, potassium sorbate)], Milk Protein, Skim Milk Powder, Stabiliser (acetylated distarch adipate, gelatin, guar gum, pectins), Yoghurt Culture, Bifidobacterium, Lactobacillus acidophilus	
Allergens	
Milk	

Processing, Manufacturing + Packing Parameters	
1. Mix and standardise the base	Butterfat = 3.5 – 3.7% Total Solids = 20.0 – 21.0
2. Homogenise:	200 Bar

Document Reference Whole Milk Summer Fruit Bio Yoghurt 100g Specification FPSPEC 001
Revision 0 1st August 2023
Owned by: Development Manager
Authorized by: Quality Manager

Page 1 of 3 383 Words English (US) 100%

Home Insert Design Layout References Mailings Review View Share

Calibri (Body) 14

AFC Artwork Approval Form

Customer:	Product:
Date Artwork received:	Reason for Origination:
Date Artwork to be checked by:	Stage:

Operations				
Criteria	✓	X	N/A	Comments
General design Layout				
Repeat Length				
Film Width				
Film repeat				
Eye mark size, position, color				
Barcode position				
Profile Coding				
Signed Operations Manager				

Sales				
Criteria	✓	X	N/A	Comments
Bar-code				
Size Descriptor				
Pack Presentation				
Price / New Flash				

Document Reference Artwork Approval Form NPD 003
Revision 0 1st August 2023
Owned by: Development Manager
Authorized by: Quality Manager

Page 1 of 3 English (US) 100%

FSQMS Record Templates

A range of Food Safety Management System, Verification and Validation Record Templates are included.

Sample FSMS Record Templates

Search

Name

- FSR 001 Management Review Record.docx
- FSR 002 Training Record.docx
- FSR CCP Validation - Metal Detection.docx
- FSR Chemical Register.docx
- FSR CIP Pipe Flow Rate Conversion Table.xlsx
- FSR CIP Programs Log.xlsx
- FSR Cleaning Schedule.docx
- FSR Complaint Investigation Form.docx
- FSR Corrective Action Request
- FSR Design and Development.docx
- FSR Dispatch and Distribution Verification Record.docx
- FSR Document Master List.docx
- FSR Drain Cleaning Procedure.docx
- FSR Engineering Hygiene Clearance Record.docx
- FSR Equipment Cleaning Procedure and Record.docx
- FSR Equipment Commissioning Checklist.docx
- FSR First Aid Dressing Issue Record.docx
- FSR Food Safety Quality System Audit Form.docx
- FSR General Cleaning Procedure.docx
- FSR GHP Audit Checklist.docx
- FSR Glass & Brittle Material Breakage Log.docx
- FSR Glass and Brittle Plastic Register.docx
- FSR Goods In Inspection Record.docx
- FSR Goods In QA Clearance Label.docx
- FSR Hygiene Policy Staff Training Record.docx
- FSR Internal Audit Corrective Action Summary.docx
- FSR Knife Breakage Report.docx
- FSR Knife Control Record.docx
- FSR Label Retention and Check
- FSR Maintenance Work Hygiene Clearance Form.docx
- FSR Metal Detection Record.docx
- FSR Non Approved Supplier Sample Plan.docx
- FSR Non Conformance Notification.docx
- FSR Non-Conformance Record.docx
- FSR Outgoing Vehicle Inspection Record.docx
- FSR Packing Traceability Record.docx
- FSR Pre Employment Medical Questionnaire.docx
- FSR Preventative Action Request
- FSR Process Change Approval Record
- FSR Process Change Minor Approval Record.docx
- FSR Process Validation Record.docx
- FSR Product Hold Label.docx
- FSR Product Recall Record.docx
- FSR Product Recall Test Record.docx
- FSR Product Recall Trace.docx
- FSR Product Release Record.docx
- FSR PRP Cleaning Verification Record.docx
- FSR QA Online Check Sheet.docx
- FSR Return to Work Form.docx
- FSR Root Cause Analysis.docx
- FSR Sample Cleaning Record.docx
- FSR Sample Equipment Cleaning Record.docx
- FSR Sample Filler Cleaning Record.docx
- FSR Shelf Life Confirmation Record.docx
- FSR Site Audit Checklist.docx
- FSR Supplier Evaluation Form.docx
- FSR Supplier Register.xlsx
- FSR Supplier Self Assessment Form.docx
- FSR Traceability Record.docx
- FSR Vehicle Hygiene Inspection Record.docx
- FSR Visitor Questionnaire.docx
- FSR Warehouse Cleaning Record.docx

Validation Records

Verification Records

Validation Records

Name

- CCP Validation - Cleaning After Nut Production.docx
- CCP Validation - Control of Brittle Materials.docx
- CCP Validation - Dispatch and Distribution Temperatures.docx
- CCP Validation - Glass Control.docx
- CCP Validation - Metal Detection.docx
- CCP Validation Cleaning and Sanitation.docx
- Prerequisite Validation - Calibration.docx
- Prerequisite Validation - Control of Visitors and Sub-Contractors.docx
- Prerequisite Validation - Dispatch and Distribution.docx
- Prerequisite Validation - Maintenance.docx
- Prerequisite Validation - Personnel Practices.docx
- Prerequisite Validation - Control of Knives.docx
- Sample Control of Foreign Matter Contamination PRP Validation.docx
- Sample Ingredients Foreign Body Control Policy Validation.docx
- Sample Personnel Hygiene and Welfare PRP Validation.docx

Verification Records

Name

- Control of Brittle Materials Verification Record.docx
- Control of First Aid Dressings Verification.docx
- Control of Knives Verification Record.docx
- Control of Visitors and Sub-Contractors Verification Record.docx
- Despatch and Distribution Verification Record.docx
- Glass & Brittle Material Breakage Procedure.docx
- Glass Policy Verification Record.docx
- Hygiene and Housekeeping Management Verification Record.docx
- Hygiene Code of Practice Verification Record.docx
- Hygiene Policy Verification Record.docx
- Ingredients Foreign Body Control Policy Verification Record.docx
- Maintenance Verification Record.docx
- Management of Cleaning Verification Record.docx
- Management of Pest Control Verification Record.docx
- Metal Detection Verification Record.docx
- Nut Handling Procedure Verification Record.docx
- Prerequisite Verification - Training.docx

FSR Label Retention and Check [Compatibility Mo...]

Home Insert Design Layout References Mailings Review View Table Design Layout Share

AFC Label Retention and Check

Date:	17/10/22	Time:	06:00 hrs	Line Number:	1	Sample:	Start Up
							
Check and Sign							
Operator 1	Anne Operator						
Operator 2	Arno Operator						
Supervisor	Sue Pervisor						

Date:	17/10/22	Time:	08:00 hrs	Line Number:	1	Sample:	Reel Change
							
Check and Sign							
Operator 1	Anne Operator						
Operator 2	Arno Operator						
Supervisor	Sue Pervisor						

Production Manager Check Date: 17/10/22 Time: 17:00hrs Sign: Paul Manager

Document Reference FSR Label Retention and Check Record
Revision 0 1st August 2022
Owned by: Technical Manager
Authorised by: General Manager

Page 1 of 1 60 Words English (US) 100%

Supplementary Internal Auditor Training & Templates

FS 2.5.4 Internal Audits & Inspections Procedure is supported by two Auditor training presentations and sample auditing forms.

The screenshot shows a Beamer presentation slide titled "Internal Audit Training". The slide features the SQFCODE logo at the top left and navigation links for "PRODUCTS" and "ABOUT" at the top right. The main content area contains the title "Internal Audit Training" and a 3D rendering of a book titled "SQF CODE EDITION 10 FOOD SAFETY MANAGEMENT SYSTEM IMPLEMENTATION PACKAGE". The left sidebar shows a list of slides, with slide 4 selected. The bottom status bar indicates "Slide 1 of 58", "English (United Kingdom)", and "Accessibility: Good to go".

The screenshot shows a Beamer presentation slide titled "Internal Auditor Training - GMP Audits". The slide features the SQFCODE logo at the top left and navigation links for "PRODUCTS" and "ABOUT" at the top right. The main content area contains the title "Internal Auditor Training - GMP Audits" and a 3D rendering of a book titled "SQF CODE EDITION 10 FOOD SAFETY MANAGEMENT SYSTEM IMPLEMENTATION PACKAGE". The left sidebar shows a list of slides, with slide 26 selected. The bottom status bar indicates "Slide 1 of 53", "English (United Kingdom)", and "Accessibility: Good to go".

Introduction to the SQF Food Safety Management System Training Modules

AutoSave SQF System Elements FSMS Guide - Read... Saved to this PC

File Home Insert Draw Design Transitions Animations Slide Show Record Review View Help

Record Share

14 SQFCODE PRODUCTS ABOUT

F5 2.1.1.1 Food Safety Policy

15 SQFCODE

2.1.1.2 Food safety culture

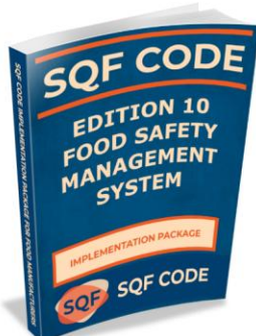
16 SQFCODE

2.1.1.2 Food safety culture

17 SQFCODE

2.1.1.2 Food safety culture

**SQF Code Edition 10 System Elements
Food Safety Management System for
Food Manufacturing**



SQF CODE

SQF CODE System Elements
Food Safety Management System for Food Manufacturing Training Guide

Slide 1 of 76 English (United Kingdom) Accessibility: Good to go

Notes

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AutoSave SQF Module 11 GMPs Guide - Read... Saved to this PC

File Home Insert Draw Design Transitions Animations Slide Show Record Review View Help

Record Share

8 SQFCODE PRODUCTS ABOUT

Module 11: Good Manufacturing Practices for Processing of Food Products

We will now go through an overview of the requirements of the SQF Code Module 11 Good Manufacturing Practices for Processing of Food Products. As we go through the presentation you will see corresponding images of documents provided with this implementation package.

9 SQFCODE

Module 11: 11.1 Site Location and Premises

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There are also some Useful Food Defense and Food Fraud Assessment Tools

FS 2.7.1A Food Defense Threat Assessment

Food Threat Assessment & Mitigation Plan Summary

Food Threat Assessment & Mitigation Plan Summary											Food Defence Mitigation Plan			
Assessment Number	Threat Category	Details	Potential Risk	Risk Assessment			Control Measures Required				Verify Controls are In Place			
				Likelihood/Vulnerability to Threat	Impact	Threat Risk Rating	Primary Control	Secondary Control	Primary Control Responsibility	Secondary Control Responsibility	Primary Control	Date	Secondary Control	Date
1	Raw Material Supply		Monitoring of Product in Market Place	3	3	9	Entrances are secured, security personnel, locks and/or alarms are installed	Ingredients are examined for possible tampering						
2	Outside Vulnerability		Outside Physical Security Measures	2	3	6	Plant boundaries are clear and secured to prevent unauthorized entry	Outside storage on the premises is protected from unauthorized access						
3	Storage		Storage Security	3	3	9	Access to storage areas is restricted	Regularly check the inventory of finished products for unexplained additions and withdrawals from existing stock.						
4	Transport		Transport Security	3	3	9	Incoming and outgoing vehicles are examined for suspicious activity	Control access to loading docks						
5	Mail Handling		Mail Handling Security	3	2	6	A food defence plan is in place	Cyber security management systems are put in place						
6	Information		Information Security	1	2	2	A food defence plan is in place	Cyber security management systems are put in place						
7	General Internal		General Internal Security Measures	1	1	1	Restricted areas are clearly identified	Ingredients are examined for possible tampering						
8	Processing Area		Processing Area Security	3	3	9								
9	Chemical/Hazardous Material Control		Chemical/Hazardous Material Control Security	3	3	9								
10	Personnel		Personnel Security Measures	3	3	9								
11	Incident Response		Incident Response			0								

FS 2.7.2A Food Fraud Assessment Template

Food Fraud Vulnerability Assessment & Plan Summary

Risks to consider are emerging and historical issues, Historical evidence of substitution or adulteration, Value of the material, Availability - e.g. a poor harvest may restrict availability and may increase the potential for adulteration, Sophistication of routine testing to identify adulterants (if testing within the supply chain is comprehensive and focused on potential fraud issues, then the likelihood is less), Country of origin, length and complexity of the supply chain

Food Fraud Vulnerability Assessment & Plan Summary											Food Fraud Mitigation Plan								
Assessment Number	Assessment Category	Details of Product or Material or Service	Details	Historical evidence of substitution or adulteration	Economic factors which may make adulteration	Ease of access to raw materials through the supply chain	Sophistication of routine testing to identify adulterants	Nature of the Raw Material	Potential Risk	Potential for Food Fraud Rating	Current Controls in Place	Risk Assessment			Control Measures Required				
												Likelihood	Public Health Consequence	Public Health Risk Rating	Primary Control	Secondary Control	Primary Control Responsibility	Secondary Control Responsibility	
1	Purchased Final Ingredient	Chocolate Topping	Supplier Barry C - India					Counterfeiting	5	Supplier Audit every 6 months	5	5	25	Raw material testing	Mass balance exercises at the raw material supplier	Supplier	Supply chain audits	Supplier	Supplier
2	Purchased Final Ingredient	Chocolate Topping	Supplier Larry B - USA					Stolen goods	3	Supplier Audit every 12 months	4	3	12	Certificates of analysis from raw material suppliers	Supply chain audits	Supplier	Supplier	Supplier	
3	Purchased Raw Ingredient	Flour for Baking	Supplier A Mills - USA					Unapproved enhancements	4	Certification to GFSI Approved Standard	5	4	20	Use of tamper evidence or seals on incoming raw materials	Enhanced supplier approval checks	Supplier	Supplier	Supplier	
4	Contract Packer	Contract Scores	Contract Pack Inc. - USA					Grey market	5	Supplier Audit every 6 months	5	5	25	Mass balance exercises at the supplier	Raw material testing	Supplier	Supplier	Supplier	
5	Purchased Contact Packaging	Cake Tray	FoodPac - Germany					Stolen goods	3	Supply to Contract Specification	3	3	9	Supply chain audits	COG with each Delivery	Supplier	Supplier	Supplier	
6	Contact Material	Detergent	Chemico Inc. - USA					No Risk	1	Supply to Contract Specification	1	1	1	Supply chain audits	COG with each Delivery	Supplier	Supplier	Supplier	
7	Purchased Non-Contact Packaging	Cardboard Box	BoxForm Inc. - USA					No Risk	1	Supply to Contract Specification	1	1	1	Certificates of analysis from raw material suppliers	Certificates of analysis from raw material suppliers	Supplier	Supplier	Supplier	
8	On-site In-Process Product	Choco Cake Mix Blend in Bulk						Stolen goods	3	Site Security	3	3	9	Certificates of analysis from raw material suppliers	Certificates of analysis from raw material suppliers	Supplier	Supplier	Supplier	
9	On-site Finished Product	Choco Cake Mix Packed						Stolen goods	3	Mass Balance exercises on site weekly	3	3	9	Certificates of analysis from raw material suppliers	Certificates of analysis from raw material suppliers	Supplier	Supplier	Supplier	
10	On-site Contact Packaging	Choco Cake Mix Bag						Counterfeiting	3	Site Security	3	3	9	Certificates of analysis from raw material suppliers	Certificates of analysis from raw material suppliers	Supplier	Supplier	Supplier	

PowerPoint Slide Show - [Food Fraud Assessment & Mitigation Plan Summary Instructions TCI]

Food Fraud Assessment & Mitigation Plan Summary Instructions

Open Excel file FS 2.7.2A Food Fraud Assessment Template

This is the main **Food Fraud Assessment Worksheet**

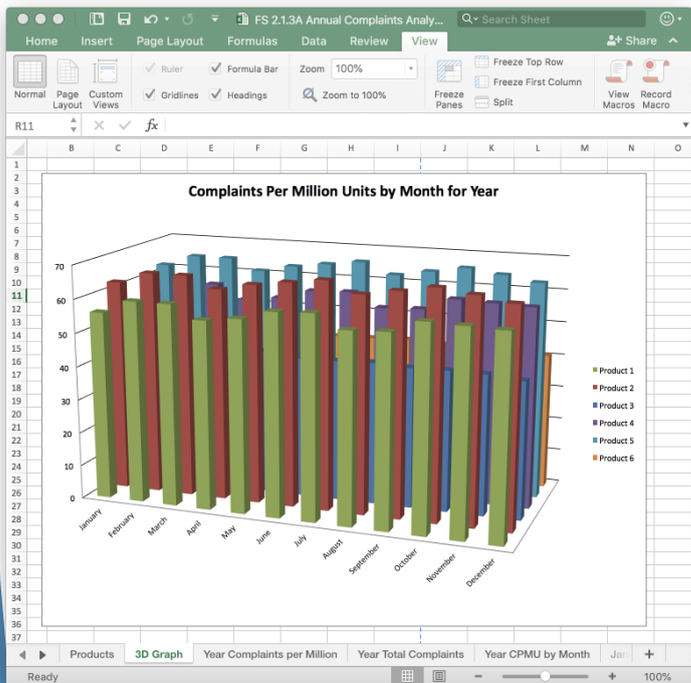
To supplement FS 2.1.3 Complaint Management Procedure, there is a Complaint Analyzer with Instruction and Guidance on Reducing Complaint levels

The image shows three pages of a Microsoft Word document titled "FS 2.1.3 Complaint Management". The pages are titled "AFC Complaint Management" and contain the following sections:

- Page 1:** Introduction, Scope, Procedure, Receipt of External Information, and Responsibilities.
- Page 2:** 5. Where found, 6. Details of any action taken by complainant, and a list of responsibilities.
- Page 3:** References, Product Recall, Corrective Action, and Complaint Investigation Form.

The image shows a presentation with two main slides:

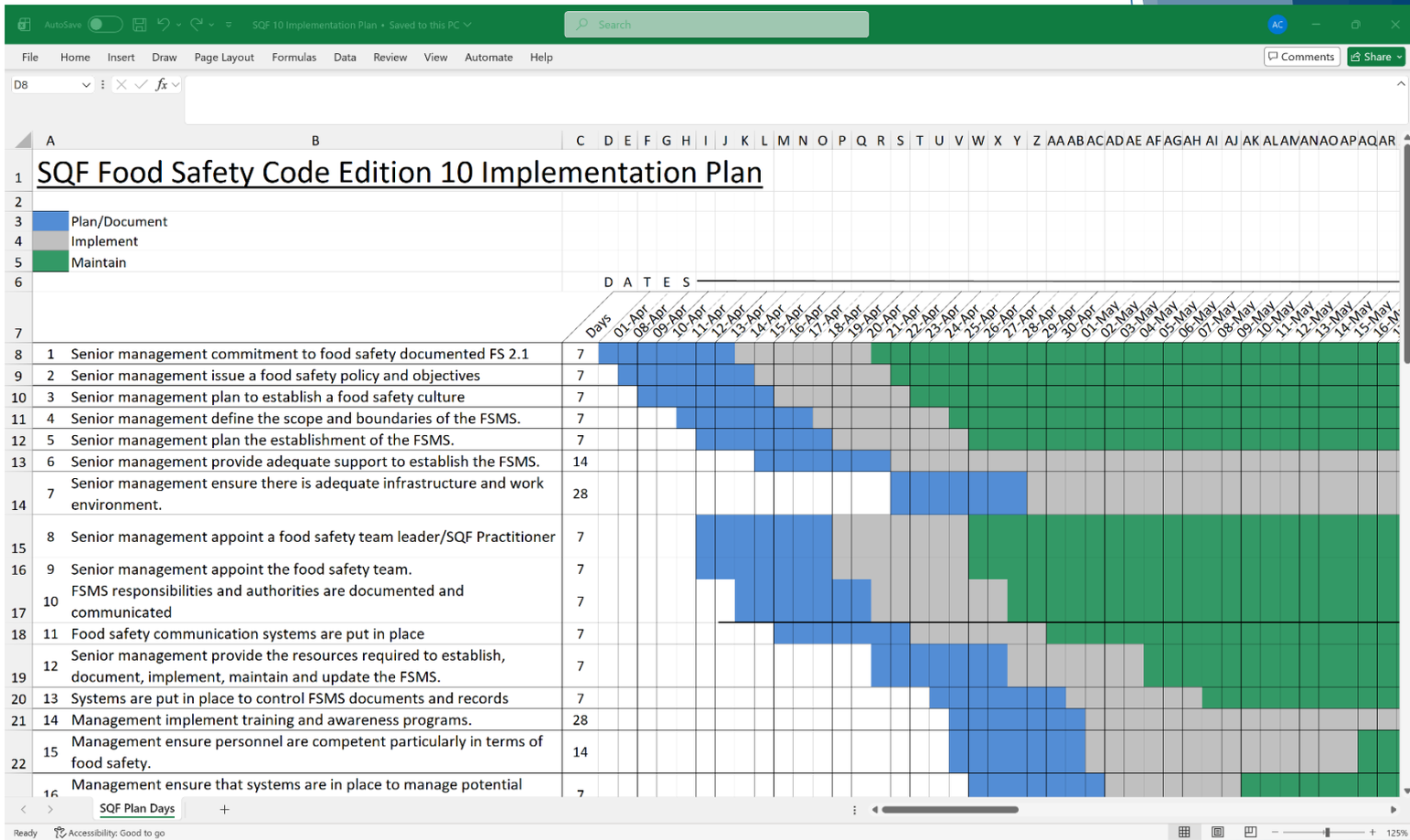
- Complaint Analyzer Instructions:** A 3D bar chart showing "Complaints Per Million Units by Month for Year" for six products (Product 1 to Product 6) across the months of the year.
- Complaint Trend Analysis:** A 3D bar chart showing "Now enter the product sales figures for the month" for six products across the months of the year.



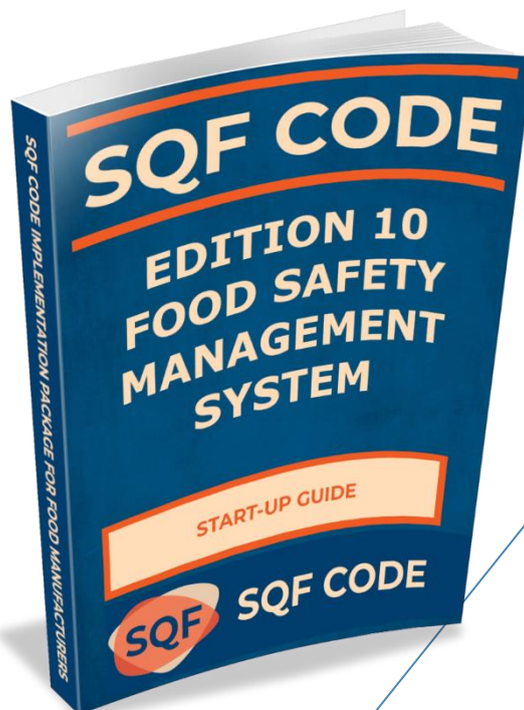
Complaint Analyzer Complaints per Million Units by Month

Month	Product					
	Product 1	Product 2	Product 3	Product 4	Product 5	Product 6
January	56	63.0	42.2	56	63.0	42.2
February	60	66.3	41.8	60	66.3	41.8
March	60	66.3	41.8	60	66.3	41.8
April	56	63.0	42.2	56	63.0	42.2
May	57.2	65.0	43.0	57.2	65.0	43.0
June	60	66.3	41.8	60	66.3	41.8
July	60.4	67.7	42.0	60.4	67.7	42.0
August	56.4	64.3	42.4	56.4	64.3	42.4
September	56.8	66.0	41.8	56.8	66.0	41.8
October	60.4	67.7	42.0	60.4	67.7	42.0
November	60	66.3	41.8	60	66.3	41.8
December	59.6	64.7	40.8	59.6	64.7	40.8

There is a SQF Implementation Plan Edition 10 which can be used to plan the development of your Food Safety Management System



Start-Up Guide - Introduction to the contents of the package



Technical Support



Free Online Technical Support

One of the unique features of our packages is that we provide technical support.

This package includes online technical support and expertise to answer your questions and assist you in developing your SQF Edition 10 Food Safety Management System until you achieve certification.

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