Facility Shutdown Management: Best Industry Practices

To Ensure a Smooth Shutdown and a Rapid Startup

by Martin Lush

Martin Lush underlines the importance of controlling plant shutdowns so that start-up is on time with minimal disruption to the ongoing supply of products.

“We all have our war stories of shutdowns dragging on for weeks, not days, and it is worrying that after many engineering interventions it can take weeks or even months before the facility performs as well as it did beforehand,” says Martin. “What can you do to make sure this important event is performed rapidly, under close control and within the cGMPs?”

Facility Shutdown Maintenance: The Context

Facility or “planned” shutdown maintenance is vital for any production facility. Access to these plants and equipment is usually restricted during routine operations, so planned shutdowns provide the opportunity for the engineering team to complete major maintenance to the plant as well as equipment.

However shutdowns can come in many shapes and sizes!

> The genuinely planned! These are usually scheduled well in advance for large-scale maintenance activities. The number of these planned shutdowns depends on the nature of the manufacturing process and how hard the plant is working, the so-called plant utilization. The greater the plant utilization, the more (preventative) maintenance is required. When companies wrongly perceive these to be a cost, not an investment…they are in trouble. This is like waiting for your car to break down rather than having it regularly serviced

> The unplanned emergency shutdown. A leaking pipe or a catastrophic equipment failure usually happens when you least expect it and usually at the worst time possible. To fix it, the plant must be shut down quickly in a controlled way

> Full and partial shutdowns. Planned shutdowns usually involve closing the entire plant. Emergency shutdowns sometimes require only partial closure, presenting some unique challenges for startup and maintaining high quality product supply

Shut downs represent a high risk to your operation and are costly!

> Lots of contractors and third parties are usually involved. All must be managed

> The build up to shutdowns can be rushed as plants frantically attempt to “catch up” with stock build and manufacturing schedules
It’s amazing how some equipment never works as well as it did before it was stopped and had parts replaced! This can lead to delays, stress and frustration as plant engineers struggle to hit deadlines for startup. This burn in period can stretch for months and lead to further interventions and costly GMP incident investigations.

- Change control! Changes to plant and equipment have to be reviewed within hours, not days or weeks; and the expertise has to be on hand to make this happen.
- Long hours. It is not uncommon for engineering teams to work 24/7 to get the job done. Fatigue + Stress = Mistakes.
- Since Job A must precede Job B, careful planning and scheduling is vital.

- Too many cooks. Contractors, engineers, validation specialists, QA, QC, Procurement, Operations; all have a part to play. Without leadership, planning, control and short interval management, the outcome could be disastrous.
- Contamination control. It is often the case that the job people least like doing is usually the most critical. In this case, post shutdown cleaning and sanitization can be a real chore yet they are critical in re-establishing environmental standards and control.

No matter what type of shutdown (planned or emergency), the key to success is organization and discipline:

- Clearly defined roles and responsibilities
- Clear handover between each activity or shutdown phase
- Attention to detail, particularly cleaning and sanitization
- Exquisite control over plant access, contractors and CHANGES to plant, process and equipment

We hope the following checklist will help you!

SHUTDOWN MANAGEMENT: THE PRACTICE

No matter what type of shutdown it is, there are usually five key phases, each with key considerations. Think of it like a relay race:

- **Phase One**: Pre-Shutdown Activities (Planning and Scheduling)
- **Phase Two**: Declassification and Handover
- **Phase Three**: Day-To-Day Management and Handover
- **Phase Four**: Reclassification and Handover
- **Phase Five**: Post-Shutdown Activities and Completion of the Shutdown Report

PHASE ONE: PRE-SHUTDOWN ACTIVITIES (PLANNING AND SCHEDULING)

- Confirm who must do what. Accountability and specific responsibility for key shutdown roles:
  - Shutdown coordinators/planners
  - Shutdown leader
  - Plant owner
- Your shutdown management team. Who must be involved:
  - Engineering
  - Validation/technical operations
  - Manufacturing operations
  - QA
  - QC
  - Procurement
  - Safety
  - Planning or S&OP team
- Generate a list of all shutdown activities to assess:
  - Resources required (who, what, when)
Timelines for each shutdown activity

Lockdown dates to prevent any further additions to the schedule that can’t be planned for

Pay particular attention to any task that will directly impact all others, e.g. power supply outage and smoke pattern testing

Contractors and third parties:

Competency screening and risk ranking. Are your contractors skilled and competent? Which will be responsible for conducting high-risk activities where the consequence of errors and mistakes could be costly?

Contractual arrangements signed and sealed. What they can and can’t do clearly defined and documented

Control, communication supervision and management: Who will check their work once completed?

Contingency planning. What if key contractors are unavailable? Do you have a plan B?

Keep an eye out for subcontractors being engaged without the knowledge and consent of the shutdown owners! They can wreak havoc

Equipment replacement, consumables and spare parts (availability and lead time). Make sure you have plenty of the obvious available:

Gowns/Tyvek suits

Cleaning agents of all types

Filters

Completion and publication of a Gantt chart showing key activities, resource requirements, roles and responsibilities for all key stakeholders and the exact support requirements:

Contractors

Manufacturing/operations

Engineering

Validation

QA

QC

Change control. Make sure that any planned changes to the plant and equipment are approved well in advance of the shutdown, not on the day

Criteria for handovers between each shutdown phase. Who will check and verify what, when and how? Making sure QA is involved in the approval process is key so that oversight can be maintained and the impact on products release assessed

It is vital to remind people that a plant in shutdown is still a GMP facility, not a building site. Make sure you have signage that gets this point across

PHASE TWO: DECLASSIFICATION AND HANDOVER

Removal of everything that can be moved from the plant. All movable equipment and materials should be bagged, tagged and stored to prevent contamination. Equipment that can’t be moved should be securely covered. Open pipes must be sealed/capped off to prevent contamination

Protection of walls and floors. During major shutdowns, walls and floors can be damaged requiring costly delays for additional repairs. If you expect a lot of heavy shutdown work, make sure you protect them

Sampling and monitoring points capped or sealed. If you have monitoring points, make sure these are sealed off

Audit (plant review) and sign-off vs. detailed checklist. Before you move to Phase Three, make sure you walk the plant to ensure that everything that should have been done has been done!

PHASE THREE: DAY-TO-DAY MANAGEMENT

Controlled access. Shutdowns are not a free invitation for anyone to enter the plant. The more people wandering around, the greater the chance
of losing control of the shutdown. Restrict access to those that matter

- Clothing requirements. Tyvek suits should be the minimum

- Contractor management:
  - Permit to work system (or equivalent) to ensure that contractor’s tasks are agreed to beforehand and verified upon completion. A central location (“control room”) for all work management is key. Visible, constantly manned…somewhere for people to go
  - Supervision. Make sure you have people providing day-to-day oversight and supervision of contractors based on the criticality (risk) of what they are doing
  - Verification upon completion
  - Remember, some contractors may not have a detailed understanding of the demands of working in a GMP environment. Make sure you provide an experienced local guide to oversee and to make sure they meet GMP requirements

- DAILY change control committees. Shutdowns can involve a lot of changes to equipment and the plant. Make sure these can be fast tracked for speedy approval with daily, face-to-face change control committee meetings for review and approval

- Requalification of equipment and plant. Any significant change to the plant and equipment usually requires requalification. Make sure it’s done

- Waste removal. Major shutdowns can lead to considerable waste. Make sure this is removed immediately and is not allowed to build up

- Cleaning. During major shutdowns make sure cleaning accompanies every major activity to ensure contamination is controlled throughout, making the final cleandown easier

- Daily “snagging.” Even the best planned shutdown rarely goes according to plan! After each day, complete a review of what is on track and what is not. What has gone according to plan…and what has not. This review of the snags or problems helps you in your contingency planning

- DAILY contingency planning with all key stakeholders. What extra resources are needed? How can additional risks be managed or contained?

- If you anticipate any delay to successful startup, inform your colleagues in planning ASAP! They hate surprises

- A DAILY “plan, do, review and adjust” must take place at the start of each day with all key stakeholders to ensure everyone knows what’s going on

**PHASE FOUR: RECLASSIFICATION AND HANDBOVER**

- Review and approval of requalification activities: Make sure that all equipment is operating within its validated state

- “Deep clean.” This is one of the most vital activities of any shutdown and yet the one activity often poorly done. Sanitizing agents applied to dirty surfaces, hard to clean locations ignored, activities rushed, insufficient contact time…can all delay efficient startup due to poor environmental control. Here are the key phases:
  - Firstly make sure your cleaning and sanitizing pattern or sequence does not contaminate already clean surfaces. Starting from the areas where product is open moving outwards; with the changing rooms and transfer hatches last on the list
  - Pre-clean environmental monitoring. It’s a good idea to take surface samples (swabs and contact plates) from a wide range of locations before any cleaning and sanitization activities. Pay attention to those notoriously difficult to clean locations
  - Detergent wash. Most sanitization agents
are only effective when applied to surfaces free from dirt. Cleaning all surfaces with a detergent is key

- Sanitization of all surfaces with a broad spectrum agent
- Sporicidal clean! Now it’s time to hit the spores with an aggressive sporicidal agent
- Post-clean down environmental monitoring. Take surface samples from locations adjacent to those sampled before the clean down activities

☐ HVAC cleanup: Shutdowns and the cleaning and sanitization activities can generate a lot of contamination and the plant requires time to “clean up” in the resting state. Good HVAC systems will remove contamination and return the plant to its “at rest” state within 24 hours

☐ Confirmation that “at rest” conditions have been achieved for non-viable particulates

☐ Plant visual inspection. One last walk-through to make sure everything is shipshape

☐ Formal handover and sign-off. This is when the plant is formally handed back to operations

PHASE FIVE: POST SHUTDOWN ACTIVITIES AND COMPLETION OF THE SHUTDOWN REPORT

☐ Confirm environmental monitoring results. The question often asked is “Should we wait for a full set of microbiological environmental results before we can start to make product?” Recognizing the costly delay this could incur forces many companies to be pragmatic and allow manufacturing to start without microbiological data to prove the environment is in control, providing that:

- Cleaning and sanitization have both been done effectively and signed off
- Monitoring for non-viable particulates confirms that the facility has returned to its resting state

- Everyone acknowledges that product manufactured during this period is done “at risk.” In other words, if the micro environmental data is out of specification, product may be rejected
- Confirm plant and equipment are working within validated parameters
- Remember, some equipment may have been reset to the OEM-recommended baseline parameters. In reality, you often operate equipment to different parameters based on operational history, performance and validation studies
- Close out any remaining change controls
- Close out any deviation incidents
- Compile lessons learned. It’s vital for everyone involved in the shutdown to do a lessons learned review before memories fade:
  - What went well?
  - What didn’t?
  - Improvements for next time?
- Complete shutdown report. It’s vital the report is completed quickly to include:
  - Executive summary: A short paragraph of what went well, what didn’t, any risks and how these have been managed or contained
  - Completed handover checklists covering all activities
  - Engineering status
  - Validation status
  - Change controls. Those closed, those open
  - Deviations. Number, type, status
  - Environmental control. Data to demonstrate the status of the plant for viable and non-viable particulates
- Gain approval of the report by the shutdown leader (the author), QA and the plant owner
CALL TO ACTION: WHAT YOU MUST DO FOR YOUR NEXT SHUTDOWN – TOP FIVE ACTION POINTS

> Customize the above and generate your own checklist for each of the key phases. Involve everyone in this process

> Make sure you have documented handover between each of the key phases. This involves the shutdown leader, the plant owner and QA reviewing completion of the key activities in the checklist

> Although good planning and coordination are key, shutdowns rarely go according to plan. Make sure your contingency plans are robust and your risk-based decision making skills are well practiced

> Remind everyone that the rules of GMP still apply: controlled access, contractor management, good documentation practices and so on

> Phase Five is key: Do the report immediately. Communicate the Executive Summary to QA (for the purpose of product release). Make sure you document lessons learned immediately while everything is fresh in your mind

ABOUT THE AUTHOR

Martin Lush has over 30 years’ experience in the pharmaceutical and healthcare industry. He has held senior management positions in QA, manufacturing, QC and supply chain auditing and has conducted audits and education programs for many hundreds of companies in over 25 countries.