Oxygen is the second leading cause for quality degradation of pharmaceuticals, water being the first. While we typically think of packaging as the primary resource used to protect drugs from the negative effects of oxygen, there are things to consider during development and manufacturing to make sure excess oxygen isn’t simply packaged or bottled with the drug product.

**Oxygen Sensitive Pharmaceuticals**

There are classes of drugs that are more susceptible to adulteration from exposure to oxygen. They include most biologics, DEA Schedule II and IV drugs, parenterals, and drugs that require an extended shelf life. Unintended oxygen exposure during production can set the conditions for quality degradation even though the drug may have passed all initial quality testing.
During research and development, formulations must be tested and characterized for long-term stability when exposed to oxygen and incorporated into the Critical Quality Attributes (CQA) for the drug product. If your oxygen exposure is a CQA there are ways to mitigate the impact and there are well-proven best practices that can be applied to manufacturing and liquid fill-finish.

Robotic Fill-Finish Lines

Believe or not, there was a time when it was common for parenteral formulations to be filled by hand. And although a manual filling method is rare today, completely isolated and automated fill-finish lines are not as common as one might think, mainly due to the high initial investment costs.

Fully robotic fill-finish capabilities in aseptic isolation units greatly reduce, even eliminate, the amount of oxygen exposure a drug is subjected to during manufacturing. This kind of fill-finish capability also reduces human contact with the pharmaceutical during production, protecting both the product and the operator. For any developer or manufacturer of oxygen sensitive pharmaceuticals, the investment in fill-finish systems that employ robotics in an aseptic environment ought to be seriously considered to ensure product quality.
**Single-Use Technology**

Modern single-use technology is another way to greatly reduce a drug product’s exposure to oxygen. State-of-the-art fill-finish lines are equipped with single-use components, but single-use bags for mixing and storage along with transfer lines can be used to reduce a product’s exposure to oxygen too. Many oxygen sensitive drugs are also highly potent, and single-use technology eliminates the need for extensive cleaning of mixers, blenders, and lines.

**Non-Destructive Testing**

Finally, using non-destructive inspection systems can test 100% of batches and include visual inspection of stopper depth, vial seal, fill levels, and detect microscopic cracks in vials, ampules, pre-filled syringes, and other primary containers. Excess headspace between a liquid product and a stopper can degrade the quality and efficacy of a medicine before it ever arrives to the patient. Likewise, lack of integrity in a stopper seal
or a hairline fracture in a prefilled syringe can mean a problem with the filling line that could be impacting each dose and can ruin an entire batch. However, if it can be detected before any of the doses leave an aseptic environment, the batch can be saved.

Summary

Excess oxygen exposure to certain classes of drugs can render them ineffective. When limited oxygen exposure is a CQA for a drug product, developers and manufacturers can take the steps to control their process to ensure quality outcomes. Robotic, isolated filling lines, single-use technology, and non-destructive testing can become the centerpiece of a quality system that ensures oxygen sensitive products arrive to patients unadulterated.

About Tom Ingallinera

Dr. Thomas Ingallinera is the Vice President of Technical Support at Pharmaceutics International, Inc (Pii) and one of Pii’s talented subject matter experts whose purpose is to solve problems. Dr. Tom has over 40 years of experience in pharmaceutical development, with an expertise in complex dosage forms. He has advised on product development programs across the continuum, from pre-formulation through commercial production. He has worked at several leading pharmaceutical companies including BioCryst Pharmaceuticals, Parenta Pharmaceuticals, AAI Pharma, Genzyme, Burroughs Welcome, and A.H. Robins. Dr. Ingallinera has a B.S. in Pharmacy with a Ph.D. in Pharmaceutics from Virginia Commonwealth University. Dr. Tom also serves as a member of the Technical Advisory Group for the Bill and Melinda Gates Foundation.
About Pii

Pharmaceutics International, Inc. (Pii) is a US-based contract development and manufacturing organization (CDMO) with a passion for solving problems efficiently with the highest quality standards.

Pii’s Hunt Valley, Maryland campus includes 70 manufacturing suites with 4 integrated aseptic filling lines delivering quality, safety, and efficiency. Our professionals have extensive experience with small and large molecule compounds, developing and manufacturing complex parenteral drugs, extended-release formulations, non-aqueous injectable drug products, and lyophilization. Learn more at https://www.pharm-int.com/