

Step by Step: Innovation & Automation

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Not every drug has to be blockbuster. And not every pharmaceutical company is necessarily interested in developing blockbusters.

There are plenty of opportunities for pharmaceutical companies to be successful by focusing their efforts on developing and producing products aimed at smaller but no less deserving patient populations.

The FDA classifies these diseases as “orphans” and even has a special approval process to fast track treatments for therapies that are needed immediately -- but might not be lucrative enough for bigger companies to develop.

This is where companies like United Therapeutics, based in Silver Spring, MD, step in and take over. The company has successfully merged cutting-edge technologies and a new facility with their desire to never let their patient population be without their medications.

A Personal Commitment

A parent would do anything for an ailing child. And the founder of United Therapeutics is no different. When Dr. Martine A. Rothblatt, the company's founder, chairman and CEO, needed to find a treatment for her daughter's Pulmonary Arterial Hypertension (PAH), she took it upon herself to research the disease. This led to acquisition of a compound that was originally developed by Burroughs Wellcome and the subsequent founding of United Therapeutics in 1996. That compound, Treprostinil, has since been formulated into the injectable Remodulin®, currently the company's flagship product. The company also markets two other products for the disease: Tyvaso®, an inhalation therapy, and Adcirca®, a solid dose treatment.

To support the production of the company's current sterile products and to meet

the needs of a growing product pipeline, the company realized the need for a flexible facility that would be able to handle a mix of aseptic liquid and lyophilized products in multiple formats.

To Go Virtual or Not

In this day and age of “virtual” pharmaceutical companies, which contract out their manufacturing operations and operate mainly as sales and marketing businesses -- it might have been easy for United Therapeutics to take this route as well. But with patients’ needs in mind and the availability of technology to allow them to do their own manufacturing -- the choice became an easy one.

Patrick Poisson, Vice-President Manufacturing, looks at the issue this way, “One of the unique things about our company, which only has about 500 employees worldwide, is what we are capable of from a manufacturing standpoint. Probably nine of ten companies out there our size would be virtual companies -- they might have some manufacturing capabilities -- but it might be just in one area. We have established commercial scale operations for small and large molecule APIs, sterile solutions, including injectables and inhalation products, and solid dose.”

He continues, “We are doing the manufacturing representative of a much larger company. Many might consider this contrary to how businesses are run today -- limit the bricks and mortar, outsource, and keep the costs down.” The choice to establish internal capabilities was a matter of adding redundancy to ultimately secure the supply chain for their products as Matthew VonEsch, Manager Manufacturing, says, “When you do full outsourcing with no second site it adds an element of risk that is certainly undesirable, especially with orphan drugs where alternatives are either limited or non-existent. By adding sterile fill capability we can now balance that risk by diverting production to the most appropriate site, internal or external, to ensure a supply is always available.”

Not Virtual but Vertical

With the decision made to construct their own facility, the next decision was where. The company’s corporate headquarters is a 70,500 sf building located in downtown Silver Spring. The new building was built on what was an empty parking lot next to the company’s original lab building which was completed in mid-2006. The challenge for the company was to build a very flexible and advanced aseptic manufacturing facility in a very “fixed” space and incorporate that space into a multi-use building.

Poisson explains, “We needed to have equipment flexibility simply because of how the facility is built. Walking up to the building, you notice there is no way to get equipment in or out post-construction. What we have is what we have. We had to be very careful and think very hard about what equipment we need to make the products that we are going to be making ten years from now or even twenty years from now. So we have equipment installed, at a significant capital investment, which we are currently not using. However we believe there is a very strong likelihood that we will. So we spent a lot of time discussing what the future might

bring.”

The Tools & Technologies

To make the facility work the way they wanted it to and to fit in the space they had available, the company turned to, and is currently using, many advanced technologies. Incorporated into their manufacturing facility is extensive use of single-use technologies, along with barrier isolation and Blow-Fill-Sealing filling lines. The reasons for using these technologies are manifold and most have to do with maintaining the flexibility of production they needed along with manufacturing their products in the space they had available.

In particular, the isolators have fulfilled a very important need for the company, as Poisson explains, “We have two filling lines making sterile products (BFS and barrier isolation) and we don’t have to worry about any ISO 5 clean rooms -- all the critical environments are self-created by the equipment.”

VonEsch continues, “Thanks to the isolators we have shrunk our footprint. If we wanted to have traditional ISO 5 areas our facility would have to be much larger, space we just didn’t have. We had a fixed space here. Basically it was this is your building – how do you make it work.”

Poisson adds “We had a greenfield facility that had very little room to work with. This was an office building -- with no manufacturing space. Permits were issued before it was decided to add manufacturing. So we had a footprint that we had to work with that there was no way to alter.”

VonEsch, “We had to go up instead of out. Traditional aseptic processing areas can take up a significant amount of space, because of the HEPA filtration around it. It’s cheaper to build facilities like that but the cost to maintain them is dramatically higher, because of the air handlers, electricity and constant gowning. Since we were space constrained, we had to make an ISO 5 area in a space that was too small for a traditional ISO 5 area.”

Recent advancements in technology have also helped the company. For example the company’s lyophilization line features auto-loading technology that wasn’t available years ago. In fact, the technology on the lyo line is so advanced and so flexible that the company can, if need be, fill a lyo product, and while its lyophilizing, turn the line around, fill a liquid product, close it, cap it, turn the line around again and then unload the lyophilizer.

The focus on implementing automation wherever it was possible was critical to the company, and equipment like the Groninger Filling line, Skan Isolators, Rommelag BFS, and Eisai automatic particle inspection machine have allowed the company to perform all manufacturing processes with a lean manufacturing team. VonEsch said, “It’s the same personnel that operate the BFS process as the liquid filling process; our manufacturing associates are becoming experts on not one advanced aseptic process, but two, and that’s another thing that makes us unique.”

VonEsch feels that this has to be way to do things in the future, "I think this has to be the trend with every manufacturer. Building a dedicated facility for a single product is limiting and not cost effective. Building a flexible facility like ours -- allows you to manufacture just about any product."

Proud Parents

Just like parents the company is justifiably proud of what they have accomplished. As Poisson points out, for him, there are a number of standouts, "I think looking back -- it's our project execution -- we completed construction in early December of 2009. So there were nails still going in until then. We were given FDA site approval for our first product on April 1, 2011, that's about fifteen and a half months to validate, manufacture stability batches, file and get approval for a commercial facility. We then followed it up with site approval of a second product on July 22, 2011. That is something, personally, that I'm very proud of. We had a focused group of people who worked hard every day to make things happen. It was a tremendous amount of work, it was daunting at times. Keep in mind this company had no prior experience running a sterile facility, so there were no resources internally to draw from. There were a lot of things that were potential roadblocks to our success that we were able to overcome. It was through the use of bringing in additional experienced people, identifying validation resources, working with our vendors, understanding our products and how we were going to make them that contributed to making this pretty big effort a success. I don't know of anyone who did what we did any faster."

The Future

With their facility running as planned, it's now time for them to look toward the future.

"We will continue to focus on operating this facility," says Poisson, "we have products in the pipeline." He continues, "And I'm excited now that we have the option to make them here. I think that's something that will benefit the company. Our timelines will be getting shorter as we can fill tomorrow if necessary, And we will be able to see the benefit in things like how quickly can we file NDA's/BLA's -- we will be able to start clinical trials quicker, all this excites me."

VonEsch offers his own views on the company's future, "Personally, I'm looking forward to developing new products and continuing to manufacture our existing commercial ones. Also, our isolator technology is new to the agency and being able to showcase that technology to the FDA and show them how we use it is a great opportunity."

"I will also continue to work with our vendors to make our equipment work even better. We are all actively involved in that process. We are all trying to learn as much as we can and take these learning's to apply to the equipment of the future. Its companies like Groninger, Skan, Rommelag, etc that want to take real world working knowledge and apply it to the equipment for tomorrow."

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Published on Chem.Info (<http://www.chem.info>)

“Perhaps the biggest takeaway is how much we accomplished with a small team. We all wanted to build a facility to make a product for our patients so they would always be assured of a supply.”

Source URL (retrieved on 07/10/2014 - 3:34am):

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