

STATE OF OUTSOURCING FACILITY SECTOR AND POSSIBILITIES FOR THE FUTURE

Meghan Murphy, PhD September 15, 2021

Background



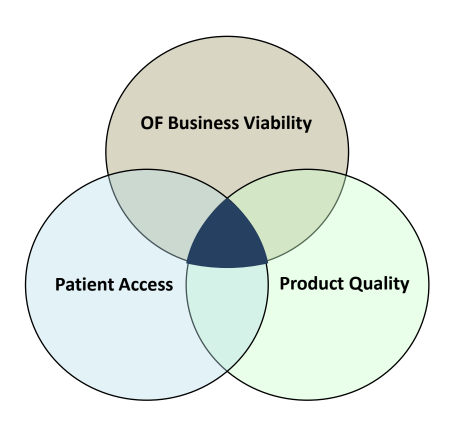
- The outsourcing facility (OF) sector has been regulated under Section 503B of the Act since 2013
- While a few firms have been operating for several years, most of the sector is comprised of relatively young companies
- As a young sector, outsourcing facilities face struggles with identity and regulatory compliance
- Outsourcing facilities serve important patient needs and this necessitates forging a clear path forward

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FDA

A Simple Framework



Sources of Information for this Presentation

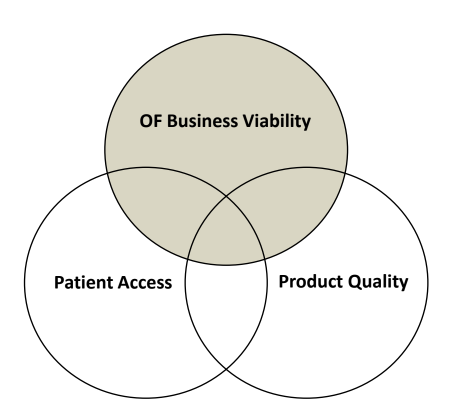


- The Compounding Quality Center of Excellence Landscape Study
 - Analysis of data
 - Annual surveys of OFs
 - Deloitte's conversations with OFs and other stakeholders

Listening sessions



OF Business Viability



The Market



76

23

\$2.3 - \$4.6B

2% - **4%**

Number of Registered
OF Facilities
(As of December 2019)

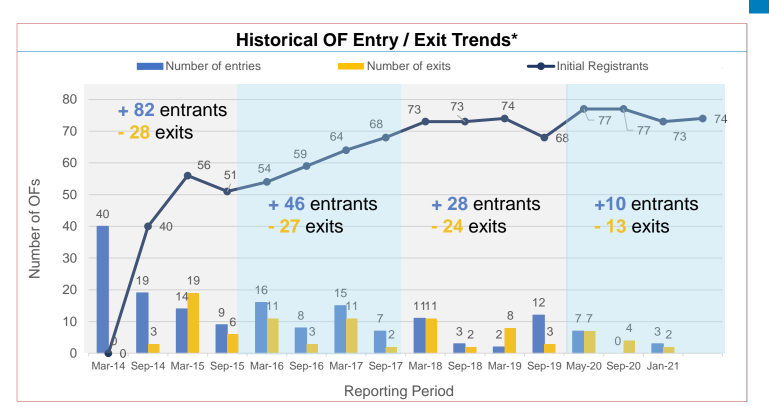
States with OFs

*Size of the 503B Market **Compounding Market Growth (503A and 503B)

*Methodology: CMS spending data indicate Medicare Part D spending on compounded drugs. Medicare Part D spending averages 30% of total prescription drug spending, which is the ratio applied to estimate total compounded drug spending. The GMI market report estimates 503B compounding holds a 52% market share. This is then applied to the total compounding market revenue to estimate the size of the 503B market; **Source: IBIS Report (2%), GMI Report (4%)

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Slowly Stabilizing, but Still in Flux



Reasons for OF Market Entry



Good fit with business model

Shortage drugs

Meet customer demand/strategic partnerships



Why Do Outsourcing Facilities Exit the Market?

Compliance issues

Difficulties in getting established

 Outsourcing facility operations not the primary business model

Operating as 503A



Top Business Challenges Identified by OFs

- Costs of maintaining and operating facilities, testing drug products, acquiring equipment, etc.
- Maintaining compliance with CGMP
- Recruiting skilled staff
- Keeping up with changing demand (either increases or inconsistencies)
- Availability of API and drug inputs



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Challenges in Identifying/Acquiring **API Supply**



- Because most OFs are not high-volume purchasers, many must depend upon brokers for API sourcing.
- Brokers vary in quality practices and reputability. Some brokers lack transparency regarding sourcing of APIs, although this appears to be improving.
- Supply chain concerns due to high overseas production and difficulties in characterizing chain of custody.
- Resource intensive to qualify suppliers, especially when multiple suppliers are needed.

rs FDA

Challenges With API Suppliers





- Competitive pricing
- Increasing demand
- Building and maintaining relationships with buyers
- Responding to drug shortages
- Using automation or technology



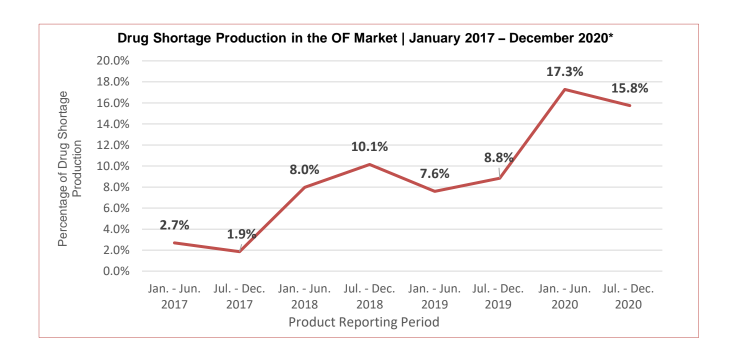
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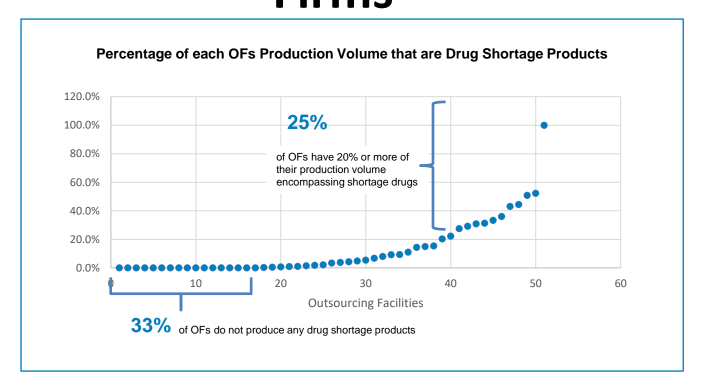






Shortage Production Varies Across Firms







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Automation Technology



 Approximately 60% of OFs claim to utilize automation technologies(based on survey data)

Most technologies center around filling and packaging/labeling

Firms not utilizing automation technology cite cost and lack of scale



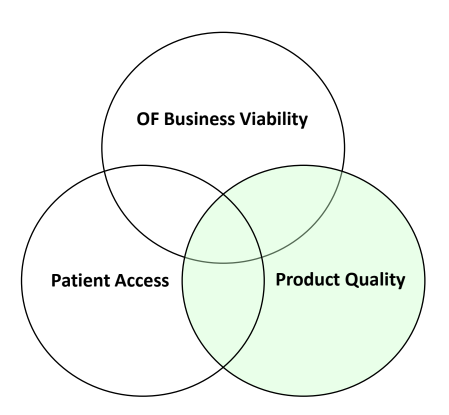


- All regulated industries are impacted by regulation—financially and otherwise.
- As a newly regulated industry, there is greater policy flux than with established industries as FDA continues to develop new policies.
- OFs are a young industry with mostly smaller companies. Firms can be more heavily influenced by policy changes due to being smaller in size.
- While public health and patient safety are always our foremost priorities, FDA is also mindful of the interplay between policy development and OF sector operations.
- OFs also cite that the differing regulatory models among states continue to be a challenge.





Product Quality





Quality—Motivating Factors

- In general, OFs are motivated to include quality practices/quality culture within their firms
- In addition to patient safety, OFs cite standardization, reliability, and customer satisfaction as motivators
- OFs also cite that purchasers tend to place more value on OFs that have more robust quality practices
- Risks of reputational harm and regulatory consequences also cited as motivators



Quality—Challenges

 Broad variability among OFs regarding quality practices and level of implementation

 Based on survey, 60% of OFs referenced utilization of SOPs to deal with quality failures, and 33% indicated use of a quality unit

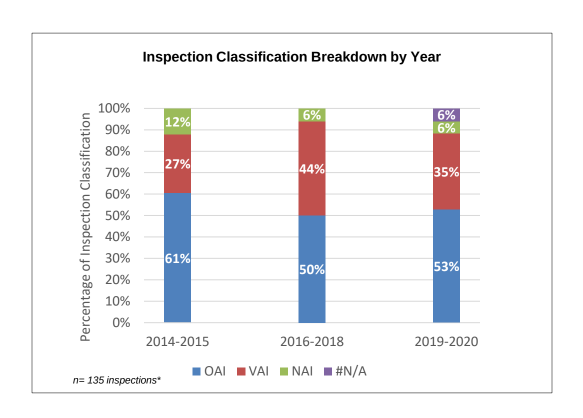


Compliance Challenges

- Compliance is an ongoing challenge, especially within the realm of CGMP
- Some OFs express a lack of clarity in reference to FDA expectations regarding 483 and Warning Letter responses
- OFs indicate FDA delays in timeliness of inspection closeouts, responses to submissions (e.g., 483 responses, Warning Letter responses)
- Some OFs indicate difficulties with understanding areas of FDA policy

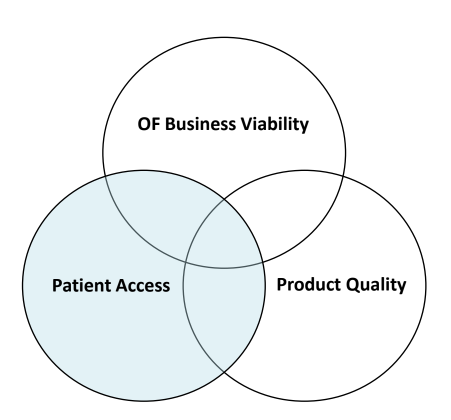


A Glimmer of Improvement



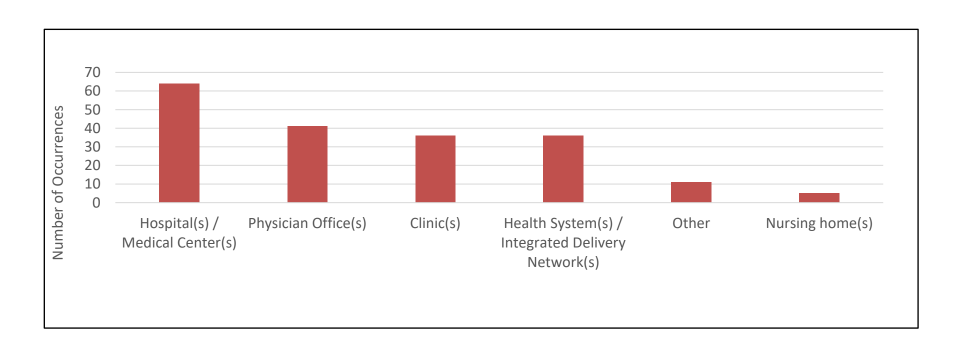


Patient Access





Who are the Customers?





Hospital Systems

 Large batch, standardized products (especially true for larger hospitals/hospital systems)

- Products that are beyond the compounding/production capabilities of in-house pharmacies
- Longer beyond use dates for products compared to inhouse or 503A pharmacy

Physician's Offices/Clinics



Smaller portfolio of specialized products

Office stock

Niche products for specialized use (ophthalmology, dermatology)





Up to 45% of OFs (per survey) work with GPOs

Typically true of larger OFs

Hospitals and health systems utilize GPOs to assist in vetting OFs

Potential Product-Specific Areas of **Demand Per Hospitals and Providers**



Shortage drugs

Ready to use formulations

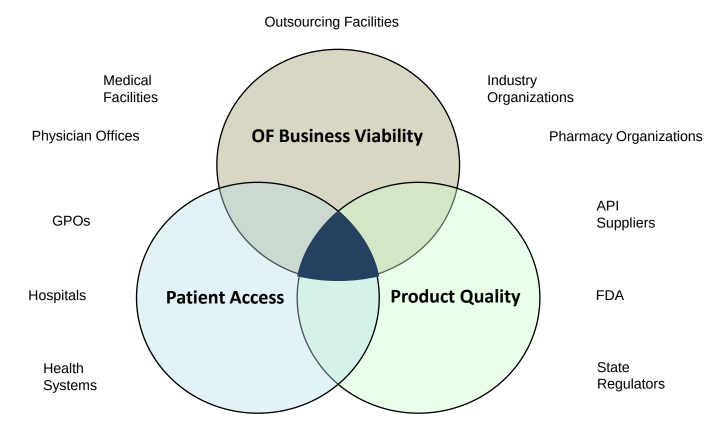
Small-batch, niche products

Key Takeaways

- OFs are young but growing and while there are difficulties, there
 are viable areas for business growth.
- Quality/compliance continues to be a pain point, but is showing signs of slight improvement and firms are motivated to improve quality.
- There is demand for OF products from diverse areas, most of which should be viable in the long-term. OFs indicate demand is growing.

The Future Involves All Stakeholders A





How Do We Get to the Center?



