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# Preparing for EU FMD & DSCSA: The Sharp Packaging Solution Perspective



If you do business in the U.S. and EU, your approach to DSCSA and FMD compliance will not be the same. Hear Sharp technical executives discuss the similarities and the critical differences between the two regulations to help you prepare.

### **VideoEuropean Union Falsified Medicines DirectiveDSCSA for ManufacturersRegulatory/ComplianceUnited States, European Union**

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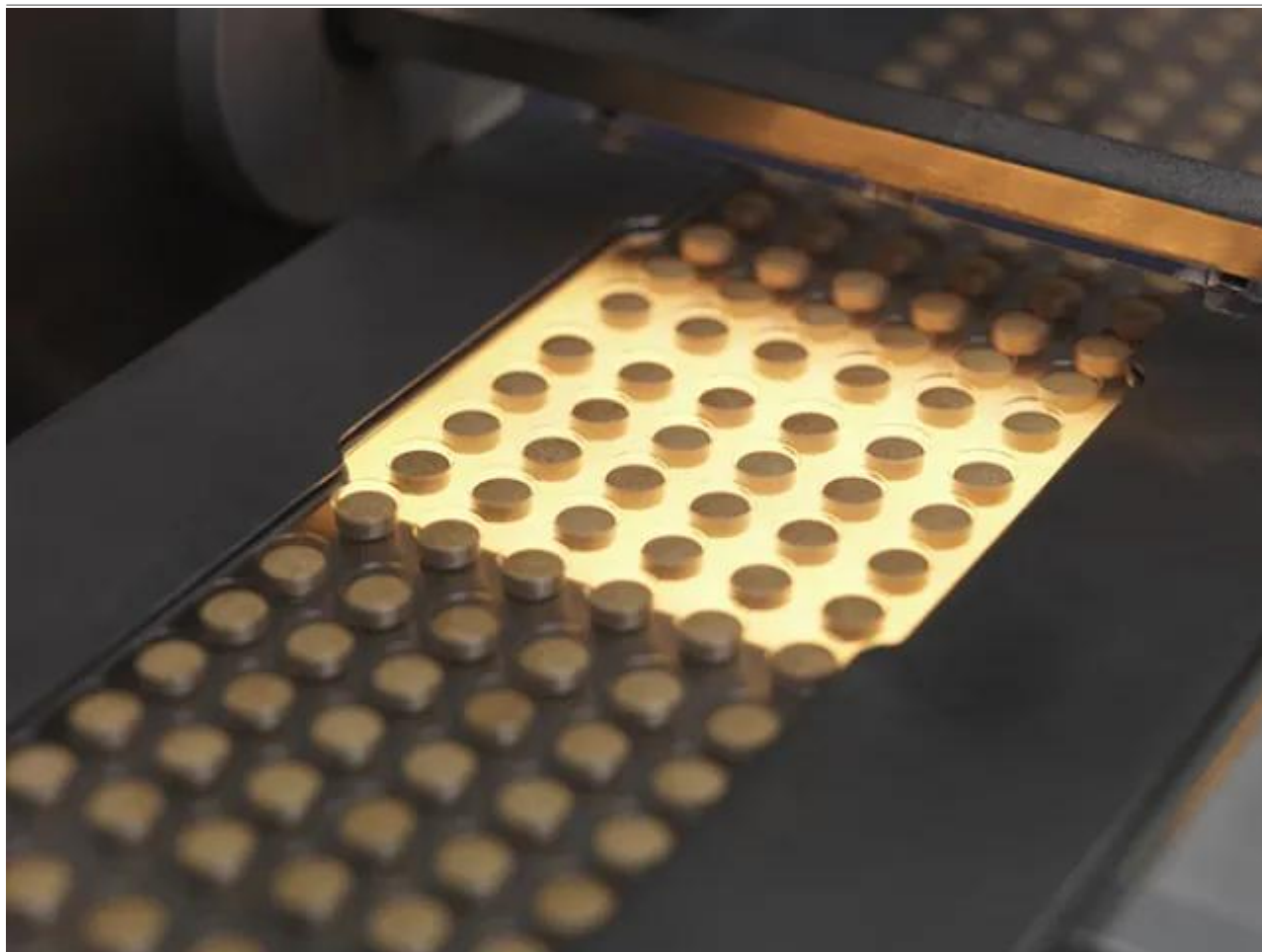
### **More Serialization and Compliance Resources**



### **Phase 3 Planning: Serialization from the Sharp Packaging Perspective**

If you're a Phase 3 pharma company, serialization will be part of your strategy. Watch Sharp SVPs discuss how they approach serialization.

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### **3M, Patheon, Reed-Lane, Sharp & Teva Share Serialization Strategies**

Watch our webinar as 3M, Patheon, Reed-Lane, Sharp and Teva share best practices to help brands and CMOs accelerate their progress to November 2017.

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## EPCIS Connection Changes post Go-Live

Lauren Catalano – Technical Services Manager



### Business Challenge & Solution

Changes to established EPCIS connections is becoming more prevalent, especially for companies like Sharp functioning in the CMO/CPO space. Technical complexities related to pathway connection changes, present the added challenge of reducing the impact to daily production activities. Allowing a smooth and unified transition to the EPCIS of choice, while working within the boundaries of business constraints is key.



#futurelink

### Case Study: Sharp Packaging Services | EPCIS Connection Changes Post Go-Live

See how Sharp Packaging Services overcame EPCIS change management challenges in the pharma supply chain with TraceLink's help.

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