

# Hybrid or Mixed Marketing Authorisation Application in the European Union

In this white paper you will learn:

- Basic EU legislation for medicinal products
- Legal basis of the application in the EU
- Hybrid applications
- How to present your Hybrid application
- Timetable for Hybrid applications under Article 10(3) of Directive 2001/83/EC
- Post authorisation
- Conclusion, Abbreviations & Bibliography