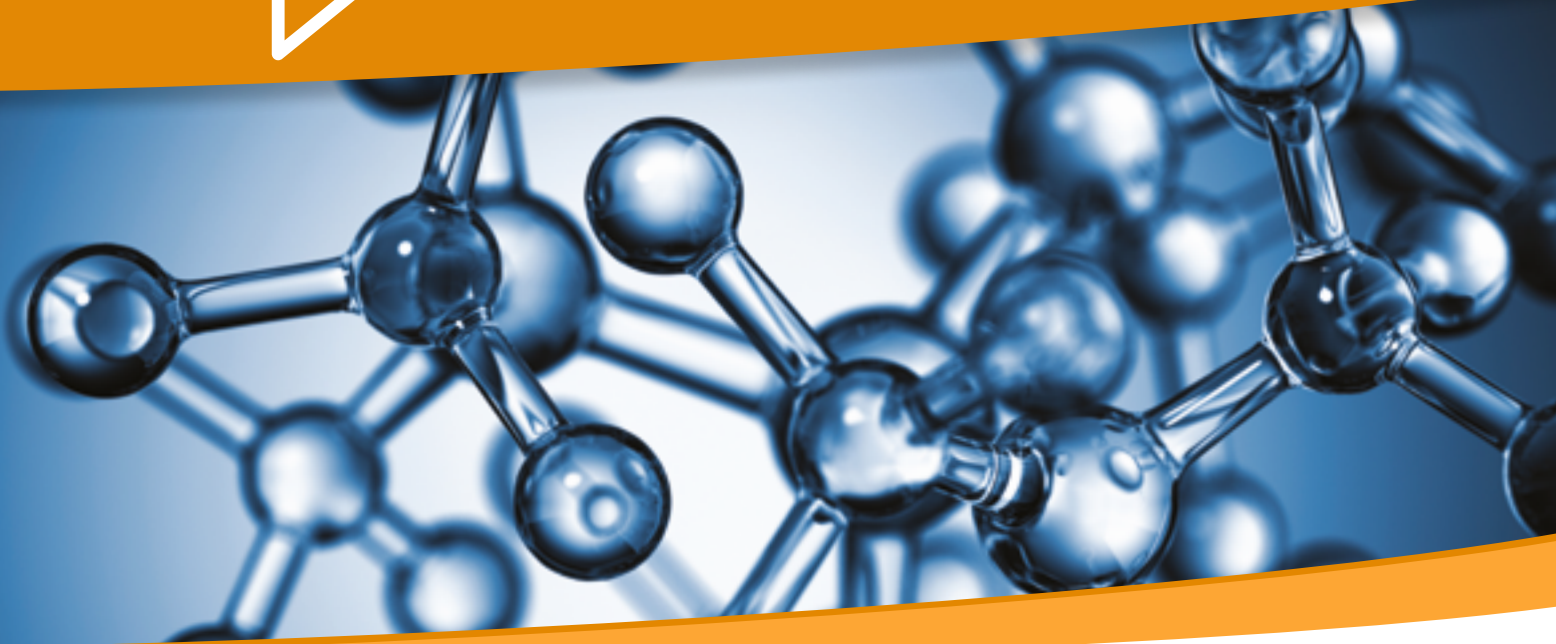






REGULATORY & CONSULTANCY



Our **Regulatory and Consultancy** team can assist you in the real obstacle course that represents obtaining a **Marketing Authorization** at the European or National level for both plant protection products and plant nutrition products (EC 1107/2009, EC 2003/2003,...).

Over 60 specialists (toxicologists, ecotoxicologists, agronomists, chemists) can help you in all areas to meet the regulatory requirements.

We work on all type of products:

- Agrochemicals
- Biopesticides 
- Adjuvants
- Fertilizers
- Biostimulants 
- Growing media

STAPHYT AGREEMENTS



Good Laboratory Practices



Good Experimentation Practices



RESEARCH TAX CREDIT
Staphyt is recognized as a Research and Development Organisation in many countries.



24 HOURS A DAY DATA,

A unique interface for monitoring the projects ensuring our customers a private and secure access to their data.

CONTACT US

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YOU NEED

- **To define the best strategy** to get the Marketing Authorization
- **To analyse available data**, identify missing data
- **To build an efficient Marketing Authorization program** that is suitable for your needs and optimized in terms of deadlines and costs
- **To manage, prepare and submit** all or part of your registration dossiers
- **To be informed about new EU and national regulations**

OUR EXPERTISE

- Recommendations about the best strategies for getting a Marketing Authorization
- Preparation of the experimentation permit
- GEP and GLP pan-European project co-ordination
- Preparation of European and national evaluation dossiers:
 - for active substances and PPP according to EC 1107/2009 regulation (all sections)
 - for nutrition products according to EC 2003/2003 or national regulations
 - for biologicals according to European and national specific requirements
- Risk assessments (Human, residues, eco-toxicology & e-fate)
- Registration dossier adaptation for Member States' requirements
- Contacts with European regulatory authorities at customer's request
- Translation (official documents, labels, SDS...)
- Regulatory watch & training

