

# Purac Biochem B.V. SITE IN Gorinchem, The Netherlands RECEIVES EXCIPACT GMP CERTIFICATION AS PHARMACEUTICAL EXCIPIENT SUPPLIER

#### Brussels, 10 June 2024

EXCIPACT is pleased to announce that the **Purac Biochem B.V. in Gorinchem in The Netherlands** has recently been awarded an EXCIPACT GMP and GDP Certificate from **SGS**, one of EXCIPACT's internationally recognised Certification Bodies.

The Certificate demonstrates that this site **Develops, manufactures and distributes of biodegradable monomers and polymers for use as pharmaceutical excipients.** according to the 2021 EXCIPACT Certification Standard for Pharmaceutical Excipient Suppliers.

Both **SGS** and their auditors had to undergo a rigorous assessment process to be EXCiPACT Registered. Both have satisfied the additional EXCiPACT competency requirements set out in the EXCiPACT annex to the ISO/IEC 17021-1:2015 standard. These include completion of the EXCiPACT Auditor Training Course, post-course written examinations and independently witnessed audits to verify their competency to the required standard. **SGS** also had to have their auditors' reports verified by their independent certification board prior to issuing the certificates.

For full details of sites currently EXCiPACT certified in 20+ countries (including: Belgium, Canada, China, France, Germany, India, Israel, Italy, Japan, The Netherlands, UK, USA, ...), see: http://www.excipact.org/certificate-holders.html.

EU and US pharmaceutical regulations require drug manufacturers to conduct either their own or thirdparty physical risk assessment audits of all of their starting material suppliers to demonstrate GMP and/or GDP compliance thus increasing the audit burden. However, using GMP and GDP standards designed specifically for pharmaceutical excipients, EXCiPACT's voluntary, independent, high quality Certification Scheme helps pharmaceutical excipient users and their suppliers to demonstrate their commitment to GMP/GDP, reducing their audit burden and costs, to assure product quality.

### Notes for the Editor

EXCIPACT provides management oversight for the voluntary, independent, high quality international EXCIPACT Certification Scheme that provides for independent 3rd party GMP and/or GDP certification of manufacturers, suppliers, and distributors of pharmaceutical excipients worldwide. The Scheme ensures patient safety through supplier quality while minimising the overall costs for assessing the excipient supply chain. It has been operational since early 2013 and there is now considerable interest among pharmaceutical excipient suppliers, customers, and regulators. For further information see <a href="https://www.excipact.org">www.excipact.org</a> or contact <a href="mailto:info@excipact.org">info@excipact.org</a>



# VWR Singapore PTE. LTD. SITE IN Singapore, Singapore RECEIVES EXCIPACT GMP CERTIFICATION AS PHARMACEUTICAL EXCIPIENT SUPPLIER

#### Brussels, 10 June 2024

EXCIPACT is pleased to announce that the **VWR Singapore PTE. LTD. in Singapore**, **Singapore** has recently been awarded an EXCIPACT GMP and GDP Certificate from **SGS**, one of EXCIPACT's internationally recognised Certification Bodies.

The Certificate demonstrates that this site **Repacks, Tests, Relabels, Stores and Distributes Dry Pharmaceutical Excipients.** according to the 2021 EXCIPACT Certification Standard for Pharmaceutical Excipient Suppliers.

Both **SGS** and their auditors had to undergo a rigorous assessment process to be EXCiPACT Registered. Both have satisfied the additional EXCiPACT competency requirements set out in the EXCiPACT annex to the ISO/IEC 17021-1:2015 standard. These include completion of the EXCiPACT Auditor Training Course, post-course written examinations and independently witnessed audits to verify their competency to the required standard. **SGS** also had to have their auditors' reports verified by their independent certification board prior to issuing the certificates.

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EU and US pharmaceutical regulations require drug manufacturers to conduct either their own or thirdparty physical risk assessment audits of all of their starting material suppliers to demonstrate GMP and/or GDP compliance thus increasing the audit burden. However, using GMP and GDP standards designed specifically for pharmaceutical excipients, EXCiPACT's voluntary, independent, high quality Certification Scheme helps pharmaceutical excipient users and their suppliers to demonstrate their commitment to GMP/GDP, reducing their audit burden and costs, to assure product quality.

### Notes for the Editor

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## Faci Metalest SLU SITE IN Zaragoza, Spain RECEIVES EXCIPACT GMP CERTIFICATION AS PHARMACEUTICAL EXCIPIENT SUPPLIER

#### Brussels, 10 June 2024

EXCIPACT is pleased to announce that the **Faci Metalest SLU in Zaragoza, Spain** has recently been awarded an EXCIPACT GMP and GDP Certificate from **SGS**, one of EXCIPACT's internationally recognised Certification Bodies.

The Certificate demonstrates that this site manufactures esters of fatty acids for use as pharmaceutical excipients according to the 2021 EXCIPACT Certification Standard for Pharmaceutical Excipient Suppliers.

Both **SGS** and their auditors had to undergo a rigorous assessment process to be EXCiPACT Registered. Both have satisfied the additional EXCiPACT competency requirements set out in the EXCiPACT annex to the ISO/IEC 17021-1:2015 standard. These include completion of the EXCiPACT Auditor Training Course, post-course written examinations and independently witnessed audits to verify their competency to the required standard. **SGS** also had to have their auditors' reports verified by their independent certification board prior to issuing the certificates.

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EU and US pharmaceutical regulations require drug manufacturers to conduct either their own or third-party physical risk assessment audits of all of their starting material suppliers to demonstrate GMP and/or GDP compliance thus increasing the audit burden. However, using GMP and GDP standards designed specifically for pharmaceutical excipients, EXCiPACT's voluntary, independent, high quality Certification Scheme helps pharmaceutical excipient users and their suppliers to demonstrate their commitment to GMP/GDP, reducing their audit burden and costs, to assure product quality.

#### Notes for the Editor

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