

# PAS 2090: PRODUCT CATEGORY RULES FOR ENVIRONMENTAL LIFE CYCLE ASSESSMENTS.

The first international standard providing a harmonised framework to assess and communicate the environmental impact of medicines across their entire lifecycle.

## PAS 2090 Development process

The standard was developed through an independent, multi-stakeholder consensus process led by BSI and sponsored by NHS England, the Office for Life Sciences (OLS) and the Pharmaceutical LCA Consortium\*.

\* Find out more here: [Link](#)

## PAS 2090 Principles

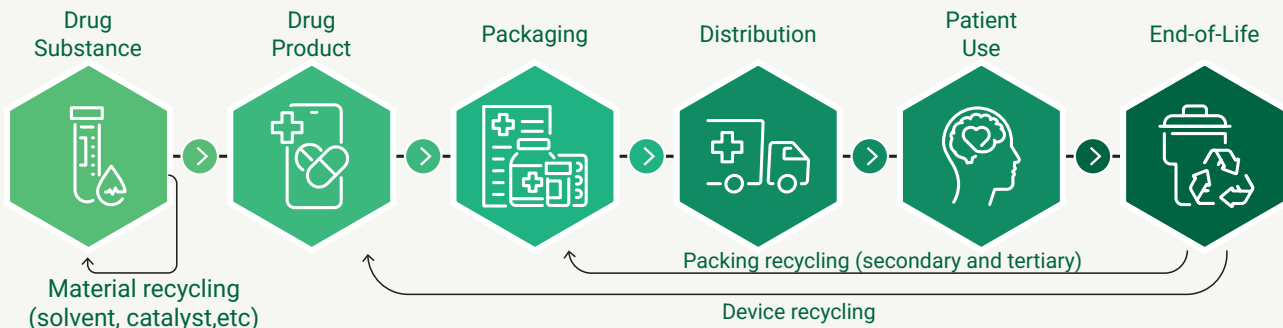
- Standardisation
- Transparency
- Collaboration

## Who is it relevant for?

-  Pharma Manufacturers
-  Healthcare Providers
-  Regulators

## KEY LIFECYCLE STAGES

PAS 2090 provides a standardised approach to assessing cradle-to-grave environmental impacts across all key stages of the product lifecycle.



## PAS 2090 STANDARD

The standard offers a globally applicable, consistent and flexible approach to LCA for pharmaceutical products.



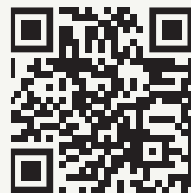
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## TECHNICAL GUIDANCE DOCUMENT

The PharmaLCA Consortium developed a technical guidance to support your PAS 2090 compliant LCA modelling.



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