Clinical Trial Transparency, Data Sharing and Disclosure Practices

Introduction

- Incyte aims to ensure that our research practices are transparent, responsible, and fully compliant with applicable laws, regulations, and guidelines.
- We have dedicated and trained staff for purposes of timely registration of clinical trials, posting of results summaries, and publication of the results in peer reviewed scientific journals.
- We support data sharing that advances science and medicine while protecting patient privacy.

Below is a summary of our policy effective as of 01May 2019. For earlier studies, different reporting requirements may have applied.

Incyte’s Clinical Trial Disclosure and Transparency Policies

Study Registration

We register Incyte-sponsored interventional and non-interventional studies that are prospectively enrolling patients on public registries such as, ClinicalTrials.gov and the European Clinical Trials Database (EUDRACT), in compliance with applicable laws and regulations.

We register expanded access programs at a product level on ClinicalTrials.gov. More information is available here.

Results Reporting on Study Registries

We post a summary of results of company-sponsored interventional and non-interventional studies that are prospectively enrolling patients, regardless of outcome on ClinicalTrials.gov and the European Clinical Trials Database (EUDRACT), in compliance with applicable laws and regulations. This also includes results summaries of clinical trials where development of the product has been discontinued.
Publication Practices

Incyte is committed to publishing high-quality publications in a responsible and ethical manner.

- Clinical studies of marketed and investigational products are submitted for publication regardless of the trial outcome.
- Publications are developed in accordance with guidelines established by the International Committee of Medical Journal Editors (www.icmje.org).

Access to Clinical Trial Data

- Incyte shares clinical trial data with external scientific and medical researchers to advance public health. Researchers may request anonymized data owned by Incyte for any study for which the product and indication are approved on or after Jan 1st 2020 in at least one major market (e.g., US, EU, JPN).
- If you are a qualified researcher interested in collaborating with Incyte and wish to request access to clinical trial data, please click here for information about the process and criteria for data sharing.