The Challenges of Writing Patient Safety Narratives for Clinical Study Reports

-Thalla Sanjeeva

Introduction

There are many different challenges that a Medical writer may encounter while writing/ managing safety narratives for Clinical Study Reports. This paper describes the importance of safety narratives and emphasizes the related challenges and gives some ideas to help you in writing/ managing patient safety narratives. The Clinical Study Report is an integrated full report of an individual clinical study of any therapeutic, prophylactic or diagnostic agent conducted in human subjects. There should be a brief narrative describing each death, each serious adverse event, and any significant adverse events that are judged to be of special interest because of clinical importance. Patient safety narratives are vital components of Clinical Study Reports (CSRs) and Pharmacovigilance (e.g. Post Marketing safety reports).

What is the Role and Purpose of Patient safety narratives?

The Clinical Study Report is an integrated full report of an individual clinical study of any therapeutic, prophylactic or diagnostic agent conducted in human subjects. There should be a brief narrative describing each death, each serious adverse event, and any significant adverse events that are judged to be of special interest because of clinical importance. Patient safety narratives are vital components of Clinical Study Reports (CSRs) and Pharmacovigilance (e.g. Post Marketing safety reports). As per International Conference on Harmonization (ICH) E3 (Section 12.3.2/14.3.3) the Narrative should describe the following:

- · The nature and intensity of the event
- · The clinical course leading up to the event
- · Timing relevant to study product administration
- Relevant Laboratory measurements
- Drug administration status
- · Postmortem findings
- Investigator's opinion, etc.

In addition, the following information should be included:

- · Subject Identifier
- · Age and sex of the subject
- · Clinical condition of the subject
- Disease being treated
- Relevant Medical history
- Concomitant and prior medication

Narratives In Clinical Study Reports

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Career Opportunities in Clinical Drug Research Rebecca Jane Anderson, 2010 It is simply amazing to me that so many of my industry coworkers stumbled upon their careers in clinical research like I did merely by chance In most cases once those opportunities were presented to us we found fulfilling and successful careers Undoubtedly other eager job seekers would also find this career path attractive If only someone would tell them about it Clinical Trials Tom Brody, 2011-11-11 Clinical Trials Study Design Endpoints and Biomarkers Drug Safety and FDA and ICH Guidelines is a practical guidebook for those engaged in clinical trial design This book details the organizations and content of clinical trials including trial design safety endpoints subgroups HRQoL consent forms and package inserts It provides extensive information on both US and international regulatory guidelines and features concrete examples of study design from the medical literature This book is intended to orient those new to clinical trial design and provide them with a better understanding of how to conduct clinical trials It will also act as a guide for the more experienced by detailing endpoint selection and illustrating how to avoid unnecessary pitfalls This book is a straightforward and valuable reference for all those involved in clinical trial design Provides extensive coverage of the study schema and related features of study design Offers a hands on reference that contains an overview of the process but more importantly details a step by step account of clinical trial design Features examples from the medical literature to highlight how investigators choose the most suitable endpoint s for clinical trial and includes graphs from real clinical trials to help explain each concept in study design Integrates clinical trial design pharmacology biochemistry cell biology and legal aspects to provide readers with a comprehensive look at all aspects of clinical trials Includes chapters on core material and important ancillary topics such as package inserts consent forms and safety reporting forms used in the United States England and Europe For complimentary access to our sample chapter chapter 24 please copy and paste this link into your browser http tinyurl com awwutvn Targeted Regulatory Writing Techniques: Clinical Documents for Drugs and Biologics Linda Fossati Wood, MaryAnn Foote, 2008-11-18 This book describes the authors standard or best practices used in writing regul ed clinical documents for the drug and biologics industry The fundamental premise of this book is that the end documents submitted to a health authority is dep dent on the beginning the planning and strategy that go into organizing written documentation Each regulatory document inherently exists within a constellation of related documents This book attempts to show the relationships between and among these documents and suggests strategies for organizing and writing these documents to maximize ef ciency while developing clear and concise text At all times and irrespective of applicable laws and guidelines good communication skills and a sense of balance are essential to adequately accurately and clearly describe a product s characteristics At no time should the reader perceive these suggestions to be the only viable solution to writing regulatory documents nor should the reader expect that these suggestions guarantee product success The audience for this book is the novice medical writer or those who would like

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