

## NACPT PHARMA COLLEGE

O/A NORTH AMERICAN COLLEGE OF PHARMACEUTICAL TECHNOLOGY www.nacptpharmacollege.com

## POST-SECONDARY DIPLOMA PHARMACEUTICAL FORMULATION & VALIDATION TECHNOLOGY



## REQUIREMENTS

Student with Minimum of Grade 12 Diploma with required science and math courses/Equivalent

> **Program Code:** PHA.FOR.VALTECH.DIP **Program Duration:** 26 Weeks of Training with Optional Job Assistance **Program**

Inquiry: 416-412-7374 Call / Text: 647-998-7374 info@nacptpharmacollege.com

# **Program Outline**

## **SEMESTER 1**

#### **GCCP01 – Good Manufacturing and Good Clinical Practices**

- The program covers concepts and requirements necessary for compliance with Good Manufacturing Practice (GMP) and Good Clinical Practice (GCP).
- GMP controls required to manufacture drug products in Canada, the United States and in many parts of the world.

#### **PMVT01 – Pharmaceutical/Bio-Pharmaceutical Manufacturing**

#### **Formulation & Validation Technology**

- Manufacturing technology, all types of process validation.
- Drug formulation.
- Manufacturing technologies.

#### **STW005 – Scientific-Technical Writing**

- Different types of technical writing.
- SOPs, Communication Methods And Issues, Validation Reports Writing, Testing Methods, CofAs, CAPA, Change Control, Gap Analysis, OOT, OOS, Deviation.

## **SEMESTER 2**

#### **ETUF05 – Collective Validation**

- Collective validation & technology regulations
- Equipment validation, Thermal validation, Utility validation, Computer & Facility Validation.

#### **PMF001 – Manufacturing Formulation 1**

- Objectives of formulation, pre-formulation techniques, manufacturing formulation techniques, and related regulations.
- Pre-formulation techniques.
- Manufacturing formulation techniques.

#### **PKV04 – Packaging Validation & Packaging Components**

- Packaging validation concept and packaging components.
- Writing protocols.
- Reports and master validation plans.
- Regulatory requirements of packaging validation.

#### **PRV03 – Process Validation**

- Process Performance Qualification And Continuous Process
  Improvements
- Process validation concept, rationale, writing protocols, reports, and master validation plans.
- Regulatory requirements of process validation.
- Focus on real work industrial scenarios.

## **SEMESTER 3**

### **PMF002 – Manufacturing Formulation 2**

- Formulation mechanism for liquid, suspension, power, fermentation, filtration, media preparation.
- Process analytical techniques and dosage evaluation.
- Effective usage liquid, suspension, powder, fermentation, filtration, media preparation, and related mechanisms in pharma, biopharmaceutical, and medical device industries.

#### **CLVM02 – Advance Cleaning Validation**

- Assessment of complex cleaning validation processes.
- The development process of new equipment/product, worst

case selection process.

• Risk-based analysis for a technical overview.

### **PPRV05 – Advance Process & Packaging Validation**

- Complex processes and packaging validation techniques.
- Risk-based analysis, technology transfer, feasibility studies.

## For more questions Visit Us at

**Toronto Campus** 9-5310 Finch Ave East Toronto, ON M1S 5E8

Brampton Campus



44 Queen St E, Brampton, ON L6V 1A2

## Contact Us

416-412-7374 | 647-998-7374 info@nacptpharmacollege.com