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08 August 2008

Dear PPMA Member:

The Seminars and Technical Committee of PPMA is very pleased with the turn out of interested participants attending our 3rd General Membership Meeting cum Technical conference which will be held in Macau from September 18 to 21, 2008.

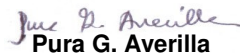
Indeed, this activity provides excellent opportunities for you to gain practical insights and knowledge as we offer this educational plant tour of the pharmaceutical manufacturing facility of **Hovione Pharma Science Ltd.**, a manufacturer of antibiotics, on September 19, 2008.

On top of this very interesting educational tour, and as committed to the general membership, we offer you an equally interesting **technical seminar** on "Case Study: **Implementing an Electronic Laboratory Information Management System in a Regulated GxP Environment**" by **Novatek International**. This topic will introduce to our members the regulatory requirements for computerized systems in the pharmaceutical industry and will explore a new system in meeting these requirements. This talk will present an actual company's implementation of an electronic laboratory information management system based on guidelines setup by US FDA, PICS and how companies can achieve and maintain cGMP using the latest system. It will also discuss the system's risk-based approach in the laboratory. Risk Management is one of the new and vital topics in the industry and which has been growing in relevance. And we are privileged to get a very knowledgeable speaker on this topic.

In this regard, the Association would like to remind you to please coordinate with our Secretariat at 893-4230 or email ppmaadel@hotmail.com. **Deadline of payment is August 22, 2008.**

Be a part of another milestone of PPMA. Thank you for your cooperation.

Very truly yours,


Pura G. Averilla

Chair, Seminars and Technical Committee

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