Introduction

Brief Overview of the Project

The Orbito project aims to deliver novel methods and a framework for rational application of predictive biopharmaceutics tools for oral drug delivery. This will be achieved through novel prospective studies to define new methodologies which will be validated using historical datasets from EFPIA partners. A combination of high quality in vitro or in silico characterizations of API and formulations will be integrated into physiologically based in silico models capturing the full complexity of oral drug absorption.

The main objectives of the project are:

- To increase understanding of the gastrointestinal drug absorption process as a prerequisite for improved biopharmaceutical prediction;
- To create new or refined in vitro and in silico methods contributing to improved in vivo predictions;
- To develop a framework for optimal use of predictive tools and preclinical models.

Project aims to establish a novel and unique database containing in vivo data of well characterized API's and oral pharmaceutical products. This database will mainly consist of novel material from the Orbito EFPIA partners.

Project database update (as of September 2nd 2013):

- Number of EFPIA companies contributed: 7
- Number of compounds uploaded to date: 61

Depending on Feasibility Populate Physiological Databases From Open Sources

The fraction of dose absorbed is affected by numerous factors which can be split into drug- and formulation-specific parameters and physiological parameters (the System). An integrated approach that considers the interplay between these factors is important for accurate prediction of the rate and extent of oral drug absorption. Physiological parameters, including, emptying rate, intestinal transit and mobility, gastro-intestinal fluid pH, secretion and reabsorption, intestinal blood flow, bile secretion, enterohepatic recirculation, and intake of food and fluids etc. form an integral component of predictive tools which rely upon extensive databases storing the mean and population variability of such parameters including, where available, their covariation. While such databases have been developed they remain significant gaps some of which will be addressed while building this database.

Some of the Areas Being Addressed

1. Circadian Variation

Many physiological processes follow circadian rhythms and the processes involved in determining the rate and extent of oral drug absorption are no exception. Gastric emptying rate and hepatic blood flow are known to be reduced significantly in the evening (night) compared to the morning (day) which may have an impact upon the rate and extent of absorption of acid-labile drugs, enteric coated formulations and drugs with significant gut-wall and hepatic metabolism. Hence, when the GI transit of drug and blood flow to liver and villi reduces at night, the rate at which drug is carried away from first-pass metabolism sites (gut-wall and liver) is lowered leading to higher first pass metabolism at night compared to the day time.

2. Viscosity of Physiologically Relevant Dissolution Systems

In order to mimic in vivo dissolution and disintegration of drugs, physiologically relevant dissolution media, mimicking gastric or intestinal juices can be employed. For such media, pH, surface tension, ionic strength & osmolality are usually considered. However, little attention has been paid to media rheology. The differential viscosity of luminal contents in the fasting and fed states can have a significant impact on dissolution and disintegration of oral dosage forms. The following components will be addressed while building this database.

- Fasted state
  - Database till date:
    - Viscosity in stomach: Chinese and Japanese
    - Viscosity in intestine: Chinese
    - Viscosity in colon: Chinese
  - Further work:
    - Viscosity in stomach: Caucasian
    - Viscosity in intestine: Japanese & Caucasian
    - Viscosity in Colon: Japanese & Caucasian

- Fed state
  - Database till date:
    - In vitro viscosity of FDA breakfast
    - In vivo viscosity of high viscosity fibres, e.g.: Dietary fibres and locust bean gum
  - Further work:
    - In vivo viscosity of FDA breakfast
    - In vivo and in vitro viscosity of high and low fat food
    - In vivo and in vitro viscosity of high and low protein food

3. Animal PBPK Models (Beagle Dog, Sprague-Dawley Rat, Generic Mouse Strain)

- Studies with preclinical species play an important role in drug development including studies with oral dosage forms. Modelling and simulation tools can, for example, be used to set appropriate dosing levels and to improve/confim understanding of drug behaviour before moving to human studies. Gastro-intestinal anatomical & physiological data for preclinical species are thus required. Data required include:
  - Small intestine and colon anatomy and physiology data (length, diameter, gastric emptying, transit time, pH, bile salt concentrations, villi morphology, unstirred boundary layer).
  - Regional small intestine, villus and colon blood flows during fasted and fed state.
  - Regional volumes of luminal fluids at steady state (influenced by secretion and reabsorption rates).
  - Regional pH and bile salt concentrations in the fasted and fed states.

4. Physiologically Relevant Buffer Systems

Development of dissolution media, already simulating in vivo physiological conditions, is expected to enhance predictions of the in vivo performance of test formulations. Bicarbonate buffer plays a major role in the in vivo gut but provides significant practical difficulties in in vitro dissolution studies. However in terms of the mechanistic modelling and simulation of dissolution in the in vivo gut it is necessary to consider the properties of bicarbonate buffer.

Population data (mean and variability) describing luminal buffer capacity and related intestinal secretion rates of relevant fluids, together with consideration of the effect of food and potentially disease state will be collected. This will then be coupled with mechanistic models for oral absorption prediction, including the modelling of surface (local) pH effects and the dissolution of pH sensitive polymers (enteric-coated formulations).