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Mathematical models for drug diffusion through the compartments of blood and tissue medium

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 Laplace transform;
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Abstract This paper is an attempt to establish the mathematical models to understand the distribution of drug administration in human body through oral and intravenous routes. Three models were formulated based on diffusion process using Fick's principle and law of mass action. The rate constants governing the law of mass action were used on the basis of the drug efficacy at different interfaces. The Laplace transform and eigenvalue methods were used to obtain the solution of the ordinary differential equations concerning the rate of change of concentration in different compartments viz. blood and tissue medium. The drug concentration in the different compartments has been computed using numerical parameters. The graphs plotted illustrate the variation of drug concentration with respect to time using MATLAB software. It has been observed from the graphs that the drug concentration decreases in the first compartment and gradually increases in other compartments.

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1. Introduction

The relation between drug intake and concentration of drug at the target site through various compartments in biological processes is considered to be the subject of great importance. The dosage and inflow and outflow of drug in the processing compartments have both favourable and adverse effects on human body. The researchers in pharmacokinetics studied the behaviour of an administered drug or chemical among various compartments of the human body over a period of time. It helps to understand the relationships between the rates of absorption, distribution and elimination process of the drug

within the body and helps to establish the desired therapeutic response. Mathematical modelling for drug diffusion constitutes an influential predictive tool to have the basic understanding of bio-transport processes. Although the mathematical modelling is theoretical in nature; however, the results established lead to realistic outcome once compared and verified empirically. In the absence of experiments, a good number of mathematical models and numerical simulations were carried out with an efficiency up to large extent. Compartment modelling plays a key role in pharmacokinetics due to local processes in each part of the compartment. Compartment model is the mathematical representation of the body or a part of the body created to study physiological or pharmacological kinetic characteristics. The body is represented as a series of compartments arranged either in series or in parallel depending upon the process or transport of material. These models can be used to understand the transport processes

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between interconnected volumes, such as the flow of drugs or any chemical within the body. A compartment model helps in understanding biological processes involved in the kinetic behaviour of a drug introduced into the body tissues. Depending upon the behaviour of drug, the body is comprising of a single or more compartment system. In one compartment model, the whole human body is considered as a homogeneous unit in which an administered drug diffuses instantaneously within the blood. In case of two compartment model, the body can be represented as two although separate but connected compartments viz. the central compartment and the peripheral one. One of the early uses of the compartment models was studied by Widmark in 1920s to model the propagation of alcohol in the body.¹

Feizabadi et al.² discussed two compartment model interacting with dynamic anti-cancer agents. Koch³ discussed the application of mathematics in pharmacokinetics by using one and two compartment models. Olga et al.⁴ developed a two-compartment mathematical model to investigate cholesterol transport in the circulatory system and its de novo synthesis in the liver. Cherrauault and Sarin⁵ studied three compartment open model with two time lags. Their model deals with the identification of exchange parameters involved in a three compartment open model with two time lags in which elimination occurs from the central compartment. Ardith and Timothy⁷ studied a mathematical model to compare bolus injection, continuous infusion for various duration, liposomal and thermoliposomal delivery of doxorubicin. Mina et al.⁸ studied the diffusion process of a drug through a skin-like membrane by making use of transformation group theoretical approach. Earlier, we⁹ studied the drug distribution in TDD systems by making use of variational finite element method taking absorption rate of drug by the tissue as decreasing function of drug concentration. Further, we used FEM to study the drug distribution in TDD systems in unsteady state case by making use of quadratic shape function.¹⁰ Moreover, Khanday and Najjar^{11,12} established the mathematical models on oxygen transport in biological tissues through capillary bed using both analytical and numerical methods. In this study, we extended the diffusion of drug in blood and tissue using three mathematical formulations and the behaviour of drug concentrations in relation with physiological parameters has been studied.

2. Mathematical model

The mathematical analysis always leads to optimal solution to various complex problems. Thus, it is imperative to establish mathematical model to estimate the drug concentration at different sites and within the blood. When the drug is orally administered, it dissolves and releases the medications into the gastrointestinal tract. The medications diffuse from there into the blood and the bloodstream takes medications to the site where it has therapeutic effect. The medications are gradually cleared from the blood by the liver and the kidneys. The flow of drugs within the body is modelled by treating the different parts of the body as compartments and then tracking the medication as it enters and leaves each compartment. The drug leaves one compartment and enters into the another one at the rate proportional to the concentration of drug present in the first compartment. The rate of drug movement between compartments is described by the first order kinetics.

The constant of proportionality is mainly determined by the drug, the compartment and general health of the individual. If $c(t)$ denotes the concentration of drug in the compartment at time t , then the rate of change of $c(t)$ is

$$\frac{dc}{dt} = \text{input rate of drug} - \text{output rate of drug.}$$

This principle is based on the law of conservation of mass and is known as the Balance Law.⁶

2.1. Model-I

A two compartment model for drug absorption and circulation through gastrointestinal tract and blood has been formulated in the beginning. The first compartment corresponds to the GI tract and from there the drug diffuses into the second compartment namely blood as shown in Fig. 1. Let $c_1(t)$ and $c_2(t)$ denote the concentration of drug in stomach or GI tract and blood stream compartments respectively. Let c_0 be the initial concentration of drug dosage. The general form of the two compartment model describing the rate of change in oral drug administration is given as

$$\left. \begin{aligned} \frac{dc_1(t)}{dt} &= -k_1 c_1(t); & c_1(0) &= c_0 \\ \frac{dc_2(t)}{dt} &= k_1 c_1(t) - k_e c_2(t); & c_2(0) &= 0 \end{aligned} \right\} \quad (1)$$

Each equation describes the change in drug concentration in their respective compartments over time. Also the quantities k_1 and $k_e (> 0)$ denote the rate constants from one compartment to another and the clearance constant. Eq. (1) are based on modelling the single dosage of drug flow via GI tract to tissue.

Solving Eq. (1), we have

$$c_1(t) = c_0 e^{-k_1 t} \quad (2)$$

$$c_2(t) = \frac{c_0 k_1}{(k_1 - k_e)} (e^{-k_e t} - e^{-k_1 t}); \quad k_1 \neq k_e \quad (3)$$

Eq. (2) corresponds to the exponential decay of drug in terms of absorption. Hence, the long term behaviour of drug concentration in GI tract will diminish to level zero.

The dosage of medicine varies from patient to patient depending upon the condition and severity of the disease. Thus, the amount of drug administration is not uniform. The oral drug medication is considered to be less efficient in some cases due to several reasons, including stomach sensitivity, liver dysfunction, delayed reaction, etc. Under such conditions, the effective and rapid drug dosage is mainly based on intravenous administration. Also in some emergency cases, medicine needed to be rapidly absorbed by the body tissues. Presence of some enzymes may break down certain delicate

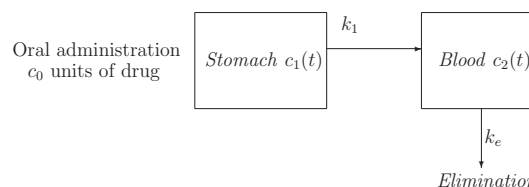


Figure 1 Simple process of drug administration through stomach and blood.

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