

Optimizing Oral Controlled Release Drug Delivery Systems using Experimental Designs

Sahilhusen I Jethara^{1,2*} and Mukesh R Patel²

¹Research scholar, Gujarat Technological University, Gujarat, India

²Shri B. M. Shah College of Pharmaceutical Education and Research, Modasa-383315, Gujarat, India

Abstract

The number of literature reports on the use of design of experiments optimization in development of drug delivery technology has been piling up steadily. This review article provides an updated bird's eye view survey account on the publications and optimization techniques of different novel controlled release delivery designs for use in oral applications. Such systematic techniques find their use in every type of conventional dosage form and novel drug delivery system. The drug delivery devices investigated for optimization using various designs include oral controlled release tablet. The present manuscript deal with various steps involved in design of experiments optimization methodology using diverse experimental designs. It also deals with a variety of showing literature findings as well as the potential application of such design of experiments procedures on optimization of assorted drug delivery systems. Such an explicit and updated review on drug delivery optimization has not been published anywhere else in the recent past.

Keywords: Controlled release; Tablet; Sustained release; Factors; Experimental designs

Abbreviations: HPMC: Hydroxypropylmethylcellulose; Na CMC: Sodium Carboxymethylcellulose; EC: Ethylcellulose; HEC: Hydroxyethylcellulose; DCP: Dicalciumphosphate; PVC: Polyvinylchloride; HPC: Hydroxypropylcellulose; MCC: Microcrystallinecellulose; PEG: Polyethylene Glycols; PVP: Polyvinylpyrrolidone

Introduction

The use of optimization techniques employing design of experiments (DoE), however, permeated the field of pharmaceutical product/process development around four decades ago. The first literature report on the rational use of optimization appeared in 1967, when a tablet of sodium salicylate was optimized using a factorial designs (FD). Since then, these systematic approaches have been put into practice in the development of drug formulations at steady pace. Despite tremendous advancements in diverse drug delivery approaches, the oral route remains the most "natural" route of drug administration. In addition, because of the low cost of oral therapy, ease of administration, and improved patient compliance associated with oral route, more than 50% of drug delivery systems available commercially are oral ones. In this context, oral controlled release drug delivery systems are quite popular, offering a number of advantages over conventional dosage forms [1,2]. Generally, the controlled release drug delivery systems for oral use are solid dosage forms, based upon the mechanism of diffusion, dissolution, or a blend of both to control the release rate of drug. These include reservoir devices wherein a polymeric membrane surrounds a drug core and matrix devices wherein the dissolved or dispersed drug is distributed uniformly in an inert polymeric matrix. Most DoE literature reports in this category are focused on optimizing the levels of these release rate-controlling polymers. DoE optimization on oral controlled release matrix delivery devices started in the early 1980s. Such devices encompass the inert matrices such as hydrophilic, hydrocolloid, silicone elastomer, and lipid matrices. The common independent variables for all of these have been the quantities of the polymers or other ingredients, while the optimized responses invariably have been the parameters characterizing in vitro dissolution profile. The other response variables

that have been optimized include disintegration time, bioavailability, and bioequivalence [3,4].

The literature reports on oral controlled release dosage forms have been compiled in various tables, categorized on the basis of various types of polymers (natural, semi-synthetic, synthetic) and the type of controlled release dosage form (matrices, dispersions, coated tablets). Table 1 depict the use of statistical experimental designs in optimization of oral sustained release (SR) matrices along with the selected drug candidate and various input variables (factors) studied.

Table 1 reports the work on DoE optimization of oral controlled release drug delivery systems, where natural, synthetic or semi-synthetic polymers have been taken as factors, invariably to control or modify the release rate of the drug. The natural polymers used comprise isapgghula husk, guar gum, xanthan gum, pectin, carrageenan, and alginic acid. Optimization reports on sustained release tablets formulated using synthetic polymers such as acrylates, poly-methacrylates, silicone elastomers, and polyethylene glycols (PEG), are shown in Table 1. Apart from polymer level, various other factors that have been optimized include tablet size, compression force, and amount of granulation liquid, lubricant, and glidants. Although most studies focused on optimizing drug release parameters, some studies involved optimization of dissolution evaluation conditions as well. Semisynthetic polymers that frequently have been employed include mainly the cellulose derivatives i.e., hydroxyl-propyl-cellulose (HPC), hydroxyl-propyl-methyl-cellulose (HPMC), hydroxyethylcellulose (HEC), sodium carboxy-methylcellulose (Sodium CMC), and ethyl-cellulose (EC). Some studies on the gums involve treatment with acid

***Corresponding author:** Sahilhusen I. Jethara, Department of Pharmaceutics, Shri B.M.Shah College of Pharmaceutical Education and Research, College Campus, India, Tel: 8460378336; E-mail: sahil.pharm4@gmail.com

Received December 17, 2014; **Accepted** December 19, 2014; **Published** January 02, 2015

Citation: Jethara SI, Patel MR (2015) Optimizing Oral Controlled Release Drug Delivery Systems using Experimental Designs. Intel Prop Rights 3: 135. doi:[10.4172/2375-4516.1000135](http://dx.doi.org/10.4172/2375-4516.1000135)

Copyright: © 2015 Jethara SI, et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

APIs	Investigated Factors	Experimental Designs	Ref.
Paracetamol	Carbopol 971P, Carbopol 71G, Tablet size	Factorial Design	[8]
Fluoride	HPMC K4M, HPMC K100 LV, Eudragit RL PO	Simplex Lattice Design	[9]
Metformin HCl	Various viscosity grades HPMC, Adhesive type, Lubricant	Response Surface Methodology	[10]
Verapamil HCl	HPMC, Na CMC	Central Composite Design	[11]
Metoprolol tartrate	HPMC, HEC, DCP	Factorial Design	[12]
Ketorolac Tromethamine	HPMC: Na CMC Ratio, EC	Factorial Design	[13]
Diltiazem HCl	Various Grades Carrageenan and Cellulose Acetate Propionate, Ionic strength, Buffer concentration	Factorial Design	[14]
Diltiazem HCl	Succinic Acid-treated Ispaghula Husk, DCP	Factorial Design	[15]
Diltiazem HCl	Guar Gum, Ispaghula Husk	Factorial Design	[16]
Diltiazem HCl	Guar Gum (Modified and Unmodified), DCP	Simplex Lattice Design	[17]
Diltiazem HCl	Modified Guar Gum, Succinic Acid, Drug content	Rotatable Central Composite Design	[18]
Diltiazem HCl	Ispaghula Husk, Water, Heating Time	Factorial Design	[19]
Ropivacaine	Pectin, EC, Binder	D-Optimal Design	[20]
Metoprolol tartrate	HPMC, Lactose: DCP ratio, magnesium Stearate, lubricant blend time, compression force	Face Centered Composite Design	[21]
Diclofenac sodium	Ispaghula husk, Lactose, MCC	Simplex Centroid Design	[22]
Diclofenac sodium	HPMC of Different Grades	Factorial Design	[23]
Diclofenac sodium	Spray-dried Rice Starch, Croscarmellose sodium, magnesium Stearate, compression force	Central Composite Design	[24]
Diclofenac sodium	EC, PVC, Talc	Factorial Design	[25]
Chlorpheniramine maleate	ψ Carrageenan, HPMC	Simplex Lattice Design	[26]
Chlorpheniramine maleate	γ-Carrageenan:cross-linked Na CMC, α-Lactose monohydrate, DCP	Simplex Mixture Design	[27]
Chlorpheniramine maleate	Na CMC, HPMC, HPC, HEC	Artificial Neural Networks, Simplex Centroid Design	[28]
Calcium phosphate	HPMC K4M, HPMC K10 M,	Simplex Mixture Design	[29]
Propranolol HCl	HPMC, Na CMC	Simplex Lattice Design, D-Optimal Design	[30]
Verapamil HCl	HPMC, Sodium Alginate	Sequential Simplex Design	[31]
Trapidil	HPMC, MCC	Central Composite Design, Artificial Neural Networks	[32]
Caffeine, ibuprofen	PEG 6000 and Acacia amount in Core	Factorial Design	[33]
Theophylline	HPC, MC, compression force	Central Composite Design	[34]
Theophylline	HPMC, HPC, MCC	Simplex Lattice Design	[35]
Theophylline	HPMC of Different Grades	Response Surface Methodology	[36]
Naftidrofuryl	Guar Gum, Xanthan Gum, MCC, Calcium Phosphate Dihydrate	Simplex Centroid Design	[37]
Naftidrofuryl	Xanthan Gum, Guar Gum	Central Composite Design	[38]
Misoprostol	HPMC, Na CMC, Lactose	Factorial Design	[39]
Alprazolam	Na CMC: Lactose ratio, HPMC 4000: HPMC 100 ratio,	Rotatable Central Composite Design	[40]
Atenolol	Various grades Carbopol	Factorial Design	[41]
Theophylline	Gelucires, Melting point, HLB, paddle rotation Speed	Factorial Design	[42]
Dextromethorphan Hydrobromide	Polydimethylsiloxane, Silicone to Silica ratio	Full Factorial Design	[43]
Potassium chloride	Silicone Elastomer Latex, PEG of Various Grades	Extreme Vertices Design	[44]
Phenyl propanolamine	Eudragit NE-40D, MCC, milling of granules before compression	Factorial Design	[45]
Ibuprofen	Eudragit (L100, RS, RSPM, RLP), EC, HPMC, HPMC phthalate	Principal Component Analysis, Response Surface Methodology	[46]
d-Chlorpheniramine maleate	EC, Eudragit, Magnesium Stearate, Talc	Factorial Design	[47]
Theophylline, Etophylline, Proxiphylline	Gelucire 50/02 & 50/13	Response Surface Methodology	[48]
Captopril	Glyceryl Monostearate, Groundnut oil	Factorial Design	[49]
Ibuprofen	Eudragit S-100, Lubricant to Glidant Ratio, Diluents, compression force	Latin Square Design	[50]
Ketoprofen	Eudragit S 100, Lactose	Box Behnken Design	[51]
Chlorpheniramine maleate	Carbopol, PVP, MCC	Extreme Vertices Design	[52]
Lobenzarit disodium	Eudragit RS-PO, MCC	Central Composite Design	[53]
Aspirin	Eudragit L100, compression force	Central Composite Design	[54]
Bumetanide	Polymer, pH Modifiers, Solubility Modifiers	Central Composite Design, D-Optimal Design	[55]
Nifedipine, Nimodipine	Carbopol 934P, Carbopol 971P, Carbopol 974P	Artificial Neural Networks	[56]
Naproxen	De-aggregating Agent, Compression Pressure	Box Behnken Design	[57]
Didanosine	Eudragit RS-PM: Ethocel 100 Ratio	Doehlert Design	[58]
Aspirin	Eudragit RS-PO, compression force	Central Composite Design	[59]
Theophylline	PEG 6000, Lactose, Stearic Acid	Response Surface Methodology	[60]

Table 1: Optimization reports on oral sustained release tablet formulations.

or alkali to modify the swelling properties of the naturally existing gum and subsequent optimization of their proportion, to be used in SR matrices. Experimental designs have also successfully been employed in the case of the core-in-cup type of compressed SR matrices, studying role of non-swell-able polymers and the other process variables in retarding the release of soluble (caffeine) and insoluble (ibuprofen) drugs employing factorial designs [5-7].

Design of experiments and optimization techniques in pharmaceutical research

The design of experiments (DOE) is an efficient procedure for planning experiments so that the data obtained can be analyzed to yield valid and objective conclusions. Experimental designs can be defined as the strategy for setting up experiments in such a manner that the information required is obtained as efficiently and precisely as possible. Well-chosen experimental designs maximize the amount of information that can be obtained for a given amount of experimental effort. Optimization of a formulation or process is finding the best possible composition or operating conditions. Determining such a composition or set of conditions is an enormous task, probably impossible and certainly unnecessary. Hence in practice, optimization may be considered as the search for a result that is satisfactory and at the same time the best possible within a limited field of search.

The purpose of optimization is to determine quantitatively the influence of the different factors together on the response variables. The number of levels is usually limited to two, but sufficient experiments are carried out to allow for interaction between factors (Figure 1) [7].

Experimental designs have long been employed to optimize various industrial products and/or processes such as;

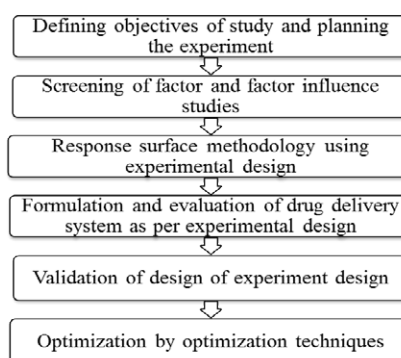
- Completely randomized designs (CRD)
- Randomized block designs (RBD)
- Screening Designs since 1946
- Simplex Lattice Design
- Latin squares designs (LSDs)
 - o Graeco-Latin squares designs
 - o Hyper-Graeco-Latin squares designs
- Factorial Designs (FDs) since 1926

- o Full factorial designs
- o Fractional factorial designs
- Plackett-Burman designs (PBDs)
- Central composite designs (CCDs) since 1951
 - o Face Centered Composite Design
 - o Rotatable Central Composite Design
- Box-Behnken designs (BBD)
- Response Surface Methodology (RSM)
- D-Optimal Design (D-OD)
- Simplex Centroid Design (SCD)
- Simplex Mixture Design (SMD) since 1958
- Sequential Simplex Design (SSDs)
- Artificial Neural Networks (ANN)
- Extreme Vertices Design (EVD)
- Doehlert Design
- Principal Component Analysis (PCA)

DOE steps:

- Problem statement
- Choice of factors, levels, and ranges
- Choice of response variable(s)
- Choice of experimental design
- Performing the experiment
- Statistical analysis
- Conclusions and recommendations

STEPS:

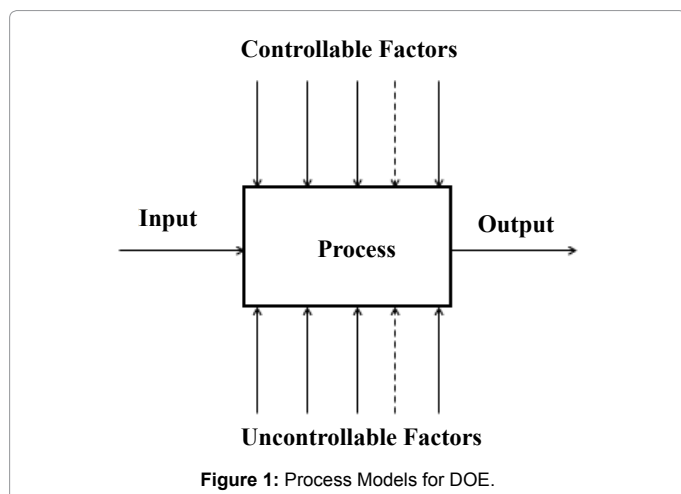


DOE applications in process development:

- Improve process yield
- Reduce variability
- Reduce development time
- Reduce overall costs

DOE objectives:

- Determine influential variables (factors)



- Determine where to set influential factors to optimize response
- Determine where to set influential factors to minimize response variability
- Determine where to set influential factors to minimize the effect of the uncontrollable factors.

DOE applications in design:

- Evaluate and compare alternatives
- Evaluate material alternatives
- Product robustness
- Determine key design parameter.

Optimizing Oral Controlled Release Tablet Formulations

An exhaustive literature search carried out by the authors in pharmaceutical journals and texts reveals that the DoE optimization techniques have been employed for almost all of these dosage forms, ranging from the simple conventional ones to that of the most intricate novel DDS. The updated literature reports unequivocally point out the increasing application of DoE techniques, with a significant shift in the focus of the formulator from optimization of the conventional formulations to that of the modern drug delivery devices [7].

Current and Future Developments

With the advent of newer, sophisticated technologies, the task of drug delivery has become more intricate, involving a greater number of resources in terms of cost, time, and energy. To circumvent these developmental hiccups, adoption of DoE analytical tools is prudently called for. Particularly, when finding the correct compromise is not straight forward, a pharmaceutical scientist should mandatorily consider the use of optimization studies.

DoE techniques have been applied with fruition on almost all kinds of drug delivery systems, not only for optimizing the formulations but their processes too. Nevertheless, there are many new drug delivery applications awaiting demonstrations. The pivotal benefits of DoE have not been thoroughly investigated in some newer drug delivery areas such as gene delivery, peptide delivery, reverse micellar systems, dendrimer based delivery systems and the like. Understanding the formulation or process variables rationally using experimental designs will help in achieving the desired goals with phenomenal ease. Experimental designs can prove to be useful, even if the primary aim is not the selection of the optimum formulation, because it tends to reveal the degree of improvement in the product characteristics as a function of the change in any excipient or process parameter(s). The major impediment in using DoE has been to envision the entire exercise as a whole. The more the formulator knows about the system, the better it can be defined, and the higher the precision with which it can be modified. The difficulties in optimizing a pharmaceutical formulation are due to the difficulty in understanding the real relationship between casual and individual responses. DoE studies can come to the rescue of the formulator, yielding much better prognostic abilities. Once the empirical relationship between the cause and the effect is unraveled, the developmental or post-developmental thoughts can be realized quite rapidly as well as rationally. Defining the relationship between the formulation or process variables and quality traits of the formulation is almost an impossible task without the application of an apt design model. Trial and error methods, in this regard, can never allow the formulator to know how close any particular formulation is to optimal

drug delivery solution. This would provide the desired impetus to the product development scientist, facilitating further evolution of research on oral controlled release drug delivery innovations and next-generation product launches.

Conclusions

The literature search indubitably ratifies the steadily increasing popularity of DoE in drug delivery optimization. Verily, the number of optimization studies would be much higher in the drug industry, where DoE methods are applied much more frequently. Because only a miniscule fraction of industrial studies is reported, most investigations remain as only in-house information. Nevertheless, the DoE usage is far from being adopted as a standard practice. Many more endeavors have to be undertaken to highlight the enormous benefits of these techniques before this can happen as a global trend. With the easy availability and affordability of DoE software, these powerful tools can be implemented with the simple click of a mouse. However, there are some key issues that depend upon the experimenter but not upon the software. These include choosing suitable responses (output variables) and factors (input variables), setting appropriate factor ranges or levels, managing the experimentation, interpreting numeric outcomes and graphic manifestations of the findings, presenting the results, and finally deciding whether to continue further with process optimization or just run confirmatory experiment(s) to validate DoE. If the experimenter has not endeavored DoE as yet or if a significant jump in information and impact in production capability has not yet been obtained, it is the most opportune time to get started. Eventually, the day will come when the benefits of DoE would be harvested by drug industry and research to their fullest advantage. Providing a relatively pithy overview, this article thus endeavors to act as a disambiguation of knowledge, and knows how to guide and provide ideas to the product development scientists in formulating varied oral controlled release drug delivery systems. I hope that my effort is going to find new application or new idea in nearer future.

Acknowledgements

The author would like to thank Dr. M. R. Patel for his comments.

References

1. Lobenberg R, Amidon GL, Vieira M (2000) Solubility as a limiting factor to drug absorption. In: Dressman JB, Lennernas H, editors. Oral Drug Absorption: Prediction and Assessment. New York: Marcel Dekker.
2. Kumar MNVR (2000) Nano and microparticles as controlled drug delivery devices. *J Pharm Pharmaceut Sci* 3: 234-258.
3. Robinson JR, Lee VHL (1987) Controlled Drug Delivery: Fundamentals and Applications. New York: Marcel Dekker.
4. Harris MR, Schwartz JB, McGinity JW (1985) Optimization of a slow release tablet formulation containing sodium sulphathiazole and a montmorillonite clay. *Drug Dev Ind Pharm* 11: 1089-1110.
5. Franz RM, Sytsma JA, Smith BP, Lucisano LJ (1987) In vitro evaluation of a mixed polymeric sustained release matrix using RSM. *J Control Release* 5: 159-172.
6. Joly F, Brossard C (1987) Development of a hydrophilic matrix for theophylline. Part 1. Physical and mechanical properties of materials and optimization of formulation. *STP Pharm. Pratiques* 3: 556-568.
7. Bhupinder S, Manju D, Vandana S, Naveen A (2005) Optimizing drug delivery systems using systematic "design of experiments." part II: Retrospect and prospects. *Crit Rev Ther Drug Car Sys* 22: 215-293.
8. Parojcic J, Duric Z, Jovanovic M, Ibric S (2004) An investigation into the factors influencing drug release from hydrophilic matrix tablets based on novel carbomer polymers. *Drug Deliv* 11: 59-65.

9. Tillotson J, Sakr A (2004) Application of multiple-response optimization to fluoride release from an extended-release matrix tablet. *Pharm Ind* 66: 601-606.
10. Li J, Dong E, Li H, Pan W (2003) Studies on the factors influencing metformin hydrochloride release from matrix tablets and the optimization of formulation. *Shenyang Yaoke Daxue Xuebao* 20: 88-92.
11. Singh B, Buhary MS, Ahuja N, Chaturvedi SC (2002) Formulation optimization and characterization of hydrophilic matrices of verapamil hydrochloride using central composite design (CCD). In: *Proceedings of 54th Indian Pharmaceutical Congress*; Pune, India.
12. Minarro M, Garcia-Montoya E, Sune-Negre JM, Tico JR (2001) Study of formulation parameters by factorial design in metoprolol tartrate matrix systems. *Drug Dev Ind Pharm* 27: 965-973.
13. Vatsaraj N, Zia H, Needham T (2002) Formulation and optimization of a sustained-release tablet of ketorolac tromethamine. *Drug Deliv* 9: 153-159.
14. Lewis GA, Chariot M (1991) Non classical experimental designs in pharmaceutical formulations. *Drug Dev Ind Pharm* 17: 1551-1570.
15. Gohel MC, Amin AF, Chhabaria MT, Panchal MK, Lalwani AN (2005) Modulation of drug release rate of diltiazem-HCl from hydrogel matrices of succinic acid-treated ispaghula husk. *Pharm Dev Technol* 5: 375-381.
16. Gohel MC, Patel MM, Amin AF (2003) Development of modified release diltiazem HCl tablets using composite index to identify optimal formulation. *Drug Dev Ind Pharm* 29: 565-574.
17. Gohel MC, Patel KV, Panchal MK, Doctor BB, Shah PD (1999) Application of simplex lattice design to the preparation of sustained-release diltiazem hydrochloride tablets using modified guar gum. *Indian J Pharm Sci* 61: 162-167.
18. Gohel MC, Panchal MK (1999) Formulation optimization of diltiazem-HCl matrix tablets containing modified guar gum using a central composite design. *Pharm Pharmacol Commun* 5: 331-338.
19. Gohel MC, Patel KV (1997) Formulation optimization of diltiazem hydrochloride matrix tablets containing modified ispaghula husk using factorial design. *Drug Dev Ind Pharm* 23: 1055-1061.
20. Ahrabi SF, Madsen G, Dyrstad K, Sande SA, Graffner C (2000) Development of pectin matrix tablets for colonic delivery of model drug ropivacaine. *Eur J Pharm Sci* 10: 43-52.
21. Rekhi GS, Nellore RV, Hussain AS, Tillman LG, Malinsowski HJ, et al., (1999) Identification of critical formulation and processing variables for metoprolol tartarate extended release (ER) matrix tablets. *J Control Release* 59: 327-342.
22. Gohel MC, Jani GK, Patel NK, Gondaliya DP (1998) Optimization of hydrophilic matrix tablet formulation of diclofenac sodium using a mixture design. *Pharm Pharmacol Commun* 4: 433-438.
23. Singh B, Gupta RK (1997) Optimization of HPMC matrix tablet formulations of diclofenac through response surface methodology. In: *Proceedings of 49th Indian Pharmaceutical Congress*; Thiruvantha puram, India.
24. Dangprasirt P, Ritthidej GC (1997) Development of diclofenac sodium controlled release solid dispersion tablet using optimization strategy. *Drug Dev Ind Pharm* 23: 843-848.
25. Pena-Romero A, Poncet M, Jinot J, Chulia D (1988) Statistical optimization of a sustained release form of sodium diclofenac on inert matrices. Part 2. Statistical optimization. *Pharm Acta Helv* 63: 333-342.
26. Bonferoni MC, Rossi S, Ferrari F, Bertoni M, Bolhuis GK, et al. (1998) On the employment of ψ carrageenan in a matrix system. Part 3. Optimization of a ψ carrageenan- HPMC hydrophilic matrix. *J Control Release* 51: 231-239.
27. Hariharan M, Gupta V, Price JC (1997) Optimization of sustained-release tablet formulations: a four component mixture experiment. *Pharm Dev Technol* 2: 365-371.
28. Hussain AS, Yu XQ, Johnson RD (1991) Application of neural computing in pharmaceutical product development. *Pharm Res* 8: 1248-1252.
29. Geoffroy JM, Fredrickson JK, Shelton JT (1998) A mixture experiment approach for controlling the dissolution rate of sustained release tablet. *Drug Dev Ind Pharm* 24: 799-806.
30. Bodea A, Leucuta SE (1997) Optimization of hydrophilic matrix tablets using a D-optimal design. *Int J Pharm* 153: 247-255.
31. Timmins P, Delargy AM, Howard JR (1997) Optimization and characterization of a pH independent extended-release hydrophilic matrix tablet. *Pharm Dev Technol* 2: 25-31.
32. Takahara J, Takayama K, Nagai T (1997) Multi-objective simultaneous optimization technique based on an artificial neural network in sustained release formulations. *J Control Release* 49: 11-20.
33. Danckwerts MP, Van der Watt JG, Moodley I (1996) The effect of processing variables on the release of ibuprofen and caffeine from controlled-release nonswellable core-in-cup compressed tablets. *Drug Dev Ind Pharm* 22: 681-687.
34. Matsumura M, Nakagami H, Yamao T, Takayama K, Nagai T (1994) Computer optimization for the formulation of controlled release theophylline tablet made of micronized low substituted hydroxypropylcellulose and methylcellulose. *Chem Pharm Bull* 42: 1902-1908.
35. Kristl A, Vojnovic D, Karba R, Mrhar A, Rubessa F (1993) Mixture design of theophylline retard formulation. *Int J Pharm* 100: 33-40.
36. Joly F, Brossard C (1987) Development of a hydrophilic matrix for theophylline. Part I Physical and mechanical properties of materials and optimization of formulation. *STP Pharm. Pratiques* 3: 556-568.
37. Waaler P, Graffner C, Muller B (1992) Optimization of a matrix tablet formulation using a mixture design. *Acta Pharm Nord* 4: 9-16.
38. Waaler PJ, Arnesen K, Graffner C, Muller BW (1992) Optimization of the amount of xanthan gum and guar gum in a matrix tablet formulation using a central composite design. *Acta Pharm Nord* 4: 291-296.
39. Shah S, Morris J, Sulaiman A, Farhadieh B, Truelove J (1992) Development of misoprostol 3 hour controlled release formulations using response surface methodology. *Drug Dev Ind Pharm* 18: 1079-1098.
40. Franz RM, Sytsma JA, Smith BP, Lucisano LJ. In vitro evaluation of a mixed polymeric sustained release matrix using RSM. *J Control Release* 5: 159-172.
41. Perez-Marcos B, Iglesias R, Gomez-Amoza JL, Martinez-Pacheco R, Souto C (1991) Mechanical and drug release properties of atenolol-carbomer hydrophilic matrix tablets. *J Control Release* 17: 267-276.
42. Ortigosa C, Guady D, Jacob M, Puech A (1991) The role of gelucire in the availability of theophylline in semisolid matrix capsules. *Pharm Acta Helv* 66: 311-315.
43. Li LC, Tu YH (1991) In vitro drug release from matrix tablets containing silicone elastomer latex. *Drug Dev Ind Pharm* 17: 2197-2214.
44. Li LC, Peck GE (1991) Water based silicone elastomer controlled release tablet film coating. V. A statistical approach. *Drug Dev Ind Pharm* 17: 27-37.
45. Meshi MS, Rivera D (1994) Factorial design of phenylpropanolamine prolonged release tablet formulations using fluid bed dryer granulator. *Drug Dev Ind Pharm* 20: 31-48.
46. Abdel-Rahman AA, Aboutaleb AE, Stamm A, Abdel-Rahman SI, Samy EM (1992) Optimization of Eudragit or cellulosic polymers ibuprofen sustained release systems using principal component analysis. *Bull Pharm Sci* 15: 63-82.
47. Fassihi R, Fabian J, Sakr AM (1996) Application of response surface methodology to design optimization in formulation of a typical controlled release system. *Drugs made in Germany* 39: 122-126.
48. Ratsimbazafy V, Brossard C (1992) Optimization of the release of theophylline derivatives from gelucire matrices. *Pharm Acta Helv* 67: 166-171.
49. Khattab IS, Abdel-rahman S, Khidr S, Abdel-Hakim O (2001) Development and optimization of sustained release captopril tablets and capsules. *Egyptian J Biotech* 10: 432-445.
50. Khan MA, Karnachi AA, Singh SK, Sastry SV, Kislalioglu SM (1995) Controlled release coprecipitates: formulation considerations. *J. Control Release* 37: 131-141.
51. Khan MA, Dib J, Reddy IK (1996) Statistical optimization of ketoprofen Eudragit S 100 coprecipitates to obtain controlled release tablets. *Drug Dev Ind Pharm* 22: 135-141.
52. Hirata M, Takayama K, Nagai T (1992) Formulation optimization of sustained release tablet of chlorpheniramine maleate by means of extreme vertices design and simultaneous optimization technique. *Chem Pharm Bull* 40: 741-746.

53. Boza A, De la Cruz Y, Jordan G, Jauregui-Haza U, Aleman A (2000) Statistical optimization of a sustained-release matrix tablet of lornoxicam. *Drug Dev Ind Pharm* 26: 1303-1307.
54. Ibric S, Jovanovic M, Duric Z, Parojcic J, Petrovic S, et al. (2003) Artificial neural networks in the modeling and optimization of aspirin extended release tablets with Eudragit L 100 as matrix substance. *AAPS Pharm SciTech* 4: 9-15.
55. Sakr A, Tillotson J (2004) Effect of extended-release polymer concentration and pH and solubility modifiers on the release of a weakly acid drug from an extended-release matrix tablet. In: Proceedings of Annual Meeting of American Association of Pharmaceutical Scientists (AAPS); Maryland, USA.
56. Sheng HL, Wang P, Tu JS, Yuan L, Pin QN (1998) Application of artificial neural networks for formulation designing of matrix tablets. *Zhongguo Yiyao Gongye Zazhi* 29: 352-354.
57. Zaghoul AA, Vaithiyalingam SR, Faltinek J, Reddy IK, Khan MA (2001) Response surface methodology to obtain naproxen controlled release tablets from its microspheres with Eudragit L100-55. *J. Microencapsul* 18: 651-662.
58. Sanchez-Lafuente C, Furlanetto S, Fernandez-Arevalo M, Alvarez-Fuentes J, Rabasco AM, et al. (2002) Didanosine extended-release matrix tablets: optimization of formulation variables using statistical experimental design. *Int J Pharm* 237: 107-118.
59. Ibric S, Jovanovic M, Duric Z, Parojcic J, Solomun L (2002) The application of generalized regression neural network in the modeling and optimization of aspirin extended release tablets with EudragitR RS PO as matrix substance. *J. Control Release* 82: 213-222.
60. Grassi M, Vojnovich D, Franceschinis E, Perissutti B (2003) Controlled release from hydrophobic matrices obtained by melt extrusion. In: Proceedings of 30th Annual Meeting of Controlled Release Society; Scotland, UK.

Citation: Jethara SI, Patel MR (2015) Optimizing Oral Controlled Release Drug Delivery Systems using Experimental Designs. Intel Prop Rights 3: 135. doi:[10.4172/2375-4516.1000135](https://doi.org/10.4172/2375-4516.1000135)

Submit your next manuscript and get advantages of OMICS Group submissions

Unique features:

- User friendly/feasible website-translation of your paper to 50 world's leading languages
- Audio Version of published paper
- Digital articles to share and explore

Special features:

- 400 Open Access Journals
- 30,000 editorial team
- 21 days rapid review process
- Quality and quick editorial, review and publication processing
- Indexing at PubMed (partial), Scopus, EBSCO, Index Copernicus and Google Scholar etc
- Sharing Option: Social Networking Enabled
- Authors, Reviewers and Editors rewarded with online Scientific Credits
- Better discount for your subsequent articles

Submit your manuscript at: <http://omicsgroup.org/editorialtracking/enzyme>

