Development of in vitro - in vivo correlations for newly optimized Nimesulide formulations

Muhammad Hanif1*, Muhammad Harris Shoaib2, Rabia Ismail Yousuf2, Farya Zafar2

1 Department of Pharmaceutics, Faculty of Pharmacy, Bahaudin Zakaria University Multan, Punjab, Pakistan, 2 Department of Pharmaceutics, Faculty of Pharmacy and Pharmaceutical Sciences, University of Karachi, Sindh, Karachi, Pakistan

* muhammad.hanif@bzu.edu.pk

Abstract

Use of the human volunteers in bioequivalence studies is being discouraged by the Food and drug administration after the introduction of biowaiver approaches. In-vitro in-vivo correlation (IVIVC) with the level A is accepted for the registration of new molecules. In the present study deconvolution technique with numeric approaches was applied after compressing and in vitro validating the 100mg Nimesulide immediate, intermediate and slow release tablets. Single centered, crossover, randomized study was conducted in four phases with a two-week washout period to obtain the plasma drug concentration data after administrating test and reference products in male healthy volunteers. Kinetica™ 4.4.1 (Thermoelectron corp, USA) was used for the calculation of two ways ANOVA with 90% CI from both log transformed and non-transformed data and Phoenix WinNonlin 7 and it’s IVIVC toolkit version 7.0 was used for the application of numeric approaches of IVIVC. Results revealed that the individual internal percentage prediction error for AUC_{inf} and C_{max} were found to be < 15% while their average values were < 10% in all medium. Numeric values of % PE at pH 6.8 and pH 7.4 (50 rpm in USP II and 100 rpm in USP I and II apparatus) were found to be (2.5842, 2.9789 and, 7.1732; 7.0944, 2.4721 and 4.350) for AUC_{inf} and (2.5842, 0.5736 and 4.6928; 5.6214, 3.0551 and -2.4711) values for C_{max} respectively. The low values of prediction errors demonstrate that the correlation model is projecting the in vivo response of each formulation. Percentage External error (% PE) was not required because individual values of percentage internal error (%PE) of C_{max} and AUC_{last} were not >15. In order to predict point to point correlation between fraction drug dissolved and drug absorbed, their mean \( r^2 \) value was found to be > 0.9112 which showed a linear correlation in slightly alkaline pH.

1. Introduction

Bioequivalence studies are considered very much important for the establishment of new generic dosage forms. It is a significant method to estimate the in vivo performance of the compound which can be used as a surrogate to determine the therapeutic efficacy [1]. Due to the extensive availability of generic compounds locally in the market, the need for bioequivalence studies are much more required as compared to the past decade [2]. Use of the human being
for the bioequivalence studies, cost of the study, subject to subject variation, unavailability of the expertise for the bioequivalence studies made them more complicated especially in case of generic product development. Pharmaceutical scientists have an urge to develop such techniques which are not only cost-effective but also useful for the establishment of generic products.

In vitro-in vivo correlations (IVIVC) plays a significant role in the product development and optimization process which is a very time consuming and costly procedure. Optimization of formulation requires modification in the composition of formulation, batch sizes and equipment and manufacturing procedures. If one or more such alterations are carried out in the formulation, then in vivo studies are required to be conducted to prove the comparison of the new formulation with a conventional product which will increase the total optimization cost and also increase the expenditure of carrying out multiple bioequivalence studies. In order to avoid these problems, IVIVC studies are carried out for the development of the pharmaceutical product [3, 4]. Several studies demonstrating the utilization of in vitro dissolution assessment as success criteria for the prediction of the bioavailability studies [5]. IVIVC not only reduces the time and also minimizes the in vivo experiments, but also recommended for regulatory purposes. Scientists reported that successful correlation can be used as a surrogate for bioequivalence studies and to support the biowaiver studies. Such studies can be useful to develop suitable dissolution specifications [6]. IVIVC is also adequate for the rationalization of therapeutically significant drug release specifications of the formulations [7].

Following FDA guidelines, four levels of IVIVC i.e. Level A, B, C and multiple C were present. Correlation level depends upon the ability of the correlation to demonstrate the plasma level profile completely, which may due to the administration of the given dosage form. Pharmacokinetic studies are considered very significant in the development of innovator dosage form but found to be the most expensive task for the pharmaceutical company. Three different types of guidelines have been established by the Food and Drug Administration (FDA) for bio-waiver studies of all types of drugs. First one is about the release pattern if the dosage forms show more than 85% release of drug within 30min it will be accepted without in vivo studies. The second one is the comparison of the prepared dosage form with the already established standard dosage form by similarity and dissimilarity factors. In vivo—in vitro (IVIVC) studies is the third type in which in vivo behavior can be predicted by using the in vitro data. BCS class II drugs categorized as a very potential candidate for IVIVC studies due to their in vitro profile as a rate-limiting step. Four levels i.e., A, B, C and multiple C of IVIVC can be developed after obtaining the in vitro and in vivo profile. Level A is very significant due to its point to point relationships of in vitro with in vivo data profile[8].

Level A showed a linear correlation and presented a point to point correlation between in vitro drug release studies and in vivo input rate. According to FDA guidelines, for the establishment of Level A correlation, formulations should be developed with altered drug release rates i.e. immediate (IR), intermediate (IntR) and slow release (SR) products or if drug release rate is condition independent then the single drug release rate is essential [9]. Level A correlation is of two-step method i.e. deconvolution step is followed by the assessment of the fraction of compound absorbed and dissolved. The detected fraction of drug absorbed is assessed by the numerical deconvolution procedure. The predicted fraction of drug absorbed is assessed using a detected fraction of drug dissolved. Now, these predicted drug absorbed values are used to estimate the predicted plasma concentrations by convolution method. The validity of the applied model is then assessed by computing the percentage prediction error (% PE) by comparing the difference between the predicted and observed values of several pharmacokinetic parameters i.e. Cmax & AUC0-∞. Sirisuth and Edington in 2002 estimated IVIVC model for naproxen and metoprolol [10]. Macha et al in 2009 reported a level A correlation for
nevirapine formulations using WinNonlin IVIVC Toolkit and found < 10% average %PE for different pharmacokinetic parameters [11].

Nimesulide is a selective COX-2 inhibitor and recommended for inflammation, pain, and fever [12]. Nimesulide is available in 100mg tablets, microcapsules and in controlled release formulations [13]. It belongs to BCS class II and showed less solubility and high permeability.

The aim of the study was to establish a Level A IVIVC to illustrate the relationship between in vitro release and in vivo behavior of different Nimesulide formulations immediate release (IR) intermediate release (InR) and slow release (SR) formulations. Phoenix WinNonlin 7 and its IVIVC toolkit version 7.0 was used for the establishment of correlation. Furthermore, the validity of the applied model was tested by computing the prediction errors.

2. Materials & methods

2.1 Materials

Nimesulide was gifted from PharmEvo Pakistan. Microcrystalline cellulose (Avicel PH102) and carboxymethylcellulose (Ac-di-sol®) were purchased from FMC, Brussels, Belgium. Octadecanoic acid, magnesium stearate, and sodium lauryl sulphate were purchased from RDH Germany. Hydroxypropyl methylcellulose (Methocel K4M) was gifted from Colorcon, USA. Potassium dihydrogen phosphate, Hydrochloric acid, Sodium hydroxide, Sodium Chloride, sodium taurocholate, citric acid, lecithin, disodium hydrogen Phosphate, Triton X 100, sodium dihydrogen phosphate, glacial acetic acid, sodium hydroxide pellets, citric acid, ammonium acetate and HPLC grade, all were purchased from Merck, Darmstadt, Germany.

2.2. Fast, medium and slow release tablets

Tablets of three different release fashions were planned by using the central composite rotatable design (CCRD). Nine formulations F-IR1, F-IR2 and F-IR3 from immediate release F-InR1, F-InR2 and F-InR3 from intermediate release and F-SR1, F-SR2, F-SR3 were from the slow release were optimized (already published) [14–16] due to their excellent preformulation results. All the formulations reported in Table 1 were in weights ranges from 400 ± 20 mg were compressed by direct compression with single punch machine under the controlled conditions of humidity and temperature.

2.3 Dissolution studies

Dissolution studies of marketed reference and compressed formulations F-IR1, F-IR2, F-IR3 from immediate release and F-InR1, F-InR2, F-InR3 from intermediate release F-SR1, F-SR2, F-SR3 were performed with USP Apparatus II at 100 rpm for 2 hours.

| Formulation | Code | Cro. Car. Na | Avicel | Mag. Ste | HPMC K4M | Tablet (mg) |
|-------------|------|--------------|--------|----------|----------|-------------|
| Immediate Release | F-IR1 | 1.00 | 60.00 | 2.00 | 1.25 | 357.00 |
| F-IR2 | 3.00 | 72.50 | 3.00 | 1.00 | 418.00 |
| F-IR3 | 1.00 | 83.00 | 1.00 | 1.00 | 444.00 |
| Intermediate Release | F-InR1 | 0.00 | 52.23 | 2.61 | 5.12 | 339.84 |
| F-InR2 | 0.00 | 54.11 | 3.25 | 14.56 | 387.68 |
| F-InR3 | 0.00 | 67.42 | 4.82 | 12.53 | 439.08 |
| Slow Release | F-SR1 | 0.00 | 34.16 | 3.61 | 31.06 | 375.32 |
| F-SR2 | 0.00 | 40.25 | 4.12 | 28.09 | 394.84 |
| F-SR3 | 0.00 | 48.25 | 3.61 | 32.15 | 435.4 |

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F-SR3 from the slow release were performed using USP I and II apparatuses having different composition of dissolution medium. The different dissolution medium used were 0.1N HCl having pH 1.2, phosphate buffer solution of pH 4.5, 6.8, 7.4, fasted state simulated gastric fluid (FaSSGF), fasted state simulated intestinal fluid (FaSSIF) and fed state simulated intestinal fluid (FeSSIF) [17–19]. Six tablets of each formulation were placed in a 900 ml dissolution medium by setting the temperature limits 37 ± 0.5°C at 100 rpm. Effect of surfactant was analyzed by using 1–3% sodium lauryl sulphate (SLS) after dissolving it in phosphate buffer of pH 7.4. An aliquot of 5 ml medium was taken out from vessels at different time intervals and an equal volume was replaced by fresh medium. Syringe filter of 45 μm was used for filtration process and drug concentration was calculated by UV spectrophotometer at 297 nm each experiment was repeated three times.

2.4 In vivo studies

In-vivo absorption studies were a single centered, crossover, randomized, in four phases with two week washout period in male healthy volunteers (age: 18–27 years) under the complete guidelines of FDA (www.fda.gov). Weight range of the volunteers was between ±10 percent of the ideal body weight. All physical examinations and medical examinations were within normal limits, Allergy history was also analyzed which was found negative. Those volunteers whose, weights and heights were not in normal range, their diagnostic tests failed in case of medical examination in a clinical situation, smokers like having more than 10 cigarettes daily and any other addiction like alcohol or volunteers on special diet user i.e., spicy, vegetarian, rich diet were excluded from the study. The study was conducted under the supervision of principal investigator and physician in a private hospital in Karachi after getting the ethical approval from Pharmacy, Ethics Committee, Bahauddin Zakariya University Multan. Volunteers were initially informed about the pros and cons of the study and a written consent was taken in this regard. For the bioequivalence study, optimized formulations from immediate release F-IR1 were compared with standard marketed brand “Nimaran®” 100mg (Bosch Pharmaceutical). Optimized intermediate F-InR2 and slow release F-SR3 formulations were used for the IVIVC considerations with deconvolution approaches. These formulations were given to subjects in a fasting condition with 250 ml of water at 8:00 am in the morning, who already having an overnight fast condition of around 10 hrs. Time of dose administered treated as t = 0 hrs. No food was administered 4 hrs postdose. Regular breakfast was given after 4 hrs of the administered dose. Five mL of blood sample was drawn at various time points i.e. 0.5, 1, 2, 4, 8, 12 and 24 hr after the administration of F-IR1 and marketed brand, Nimaran. For intermediate release formulation, blood sampling time was found to be i.e. 0.5, 1, 2, 4, 8, 12, 24, 36 and 48 hr. For slow release formulation, sampling time was found to be 1, 2, 4, 8, 12, 24, 36 and 72 hr. Plasma was separated using centrifuge machine at 4000 rpm for 10 minutes. Samples were frozen at—20°C. Frozen samples were thawed and plasma drug concentrations were estimated after analyzing them using already validated HPLC the method (work already reported) [20].

2.5 In vitro data analysis

Two types of dissolution testers i.e., USP dissolution apparatus I (basket method) and USP dissolution apparatus II (Paddle method) (DT 600, Erweka, Heusenstamm, Germany) were used for in-vitro drug release studies. Twelve tablets of each formulation Reference brand (Nimaran), F-IR1, F-InR2, F-SR3 were placed in 900 ml dissolution medium by setting the temperature limits 37 ± 0.5°C at two different speeds 50 and 100 rpm in both dissolution apparatuses. Five mL aliquot was drawn from dissolution mediums having Nimaran tablets and
F-IR1 formulation at 0.5, 1, 2, 4, 8, 12 hr. Similarly, the same volume of sample was taken for F-InR2 with the additional sampling time points of 24, 36 and 48 hrs. The concentration of Nimesulide from F-SR3 was observed after taking 5 mL of dissolution medium at 1, 2, 4, 8, 12, 24, 36 and 72 hr. The similarity between marketed and compressed immediate release formulation F-IR1 was calculated by similarity factor \( f_2 \), applied by the following equation.

\[
\text{\(f_2\) = \(50 \times \log \left\{ \left[ 1 + \frac{1}{n} \sum_{j=1}^{n} |R_j - T_j|^2 \right]^{-0.5} \times 100 \right\}\)
\]

Where \( n \) is the Number of samples, \( R_j \) and \( T_j \) are the Percentage release of reference and tests brands at different times respectively. Two formulations should be considered as similar if the \( f_2 \) value is more than 50.

### 2.6 In vivo data analysis

Pharmacokinetic parameters of reference (marketed research brand) and compressed F-IR1, F-InR2 and F-SR3 were calculated using Kinetica\textsuperscript{TM} 4.4.1 (Thermoelectron, USA). Data was successfully fitted into oral two-compartment model and various other compartmental and non-compartmental parameters were calculated. Similarly, bioequivalence of F-IR1 and innovator products were carried out. Different bioequivalence attributes i.e., \( \text{AUC}_{\text{last}} \), \( \text{AUC}_{0-\infty} \), \( \text{AUC}_{\text{tot}} \), \( \text{C}_{\text{max,calc}} \) and \( \text{T}_{\text{max,calc}} \). Of both reference and test products were assessed using two-way ANOVA methods. Schirmann’s two-one sided \( t \)-test was applied for the verification of the bioequivalence assessment. Products bioequivalence were established by using 90\% confidence interval (CI) values for reference and test products and the ratio of the values was targeted in the range of 0.8–1.25 for log-transformed and 0.8–1.20 for non-log transformed data respectively. For non-parametric assessment and carry-over effect, statistical software SPSS 20.0 (SPSS Inc.) was used.

### 2.7 Establishing in vitro- in vivo correlation (IVIVC)

Phoenix WinNonlin 7 and it’s IVIVC toolkit version 7.0 (Certara USA, Inc., 100 Overlook Center, Suite 101, Princeton, NJ, USA) was used to estimate the absorption of Nimesulide. In-vitro release evaluation of Nimesulide was conducted using the Weibull model and curve shape was evaluated by shape factor \( \beta \). Implicit, numeric and analytical are three different deconvolution method approaches used in previous literature among them numeric method with point area was used for the IVIVC studies. For the establishment IVIVC of level A the percentage prediction error of \( \text{C}_{\text{max}} \) and AUC was calculated.

\[
\text{C}(t) = \sum_{n=0}^{n=\text{max}} f(t_n) \text{C}_{\delta}(t - t_n) \text{d}t
\]

Where, \( f(t) \): Input rate; \( \text{C}(t) \): Response; \( \delta \): Unit impulse; \( \text{C}_{\delta}(t) \): Unit impulse response; \( \text{C}_{\delta}(t) \): Impulse response (Khan et al., 2015).

### 2.8 Predictability error for IVIVC studies (Level A)

Internal (%PE) and external (%PE) prediction errors were used in the determination of IVIVC. Internal prediction for individual formulation was estimated by its \( \text{C}_{\text{max}} \) and AUC values. Prediction error was used to assess the comparison between observed and predicted bioavailability. In the present study, external prediction error was not used because the values of internal \% PE of AUC and \( \text{C}_{\text{max}} \) were within the adequate limits. For the IVIVC predictability, the accepted limits for \%PE for \( \text{C}_{\text{max}} \) and AUC were \( \leq 10 \) [9]. The %PE\text{AUC} and % PE\text{C}_{\text{max}} can
be calculated using the following equations:

\[
\% PE_{AUC} = \frac{(AUC_{observed} - AUC_{predicted})}{AUC_{observed}} \times 100
\]

\[
\% PE_{Cmax} = \frac{(C_{max,observed} - C_{max,predicted})}{C_{max,observed}} \times 100
\]

3. Results and discussion

3.1 In vitro dissolution studies

More than 85% of Nimesulide was released from F-IR1, F-IR2 and F-IR3 within 60 minutes. pH-dependent percentage of Nimesulide released was observed in the different dissolution medium. Comparatively less amount of drug release was observed at pH 1.2 and 4.5 medium, which was may be due to weak acidic nature and pKa values of HPMC K4M. It was increased pH 6.8 and 7.4, may be due to its dissociation and micelle formation after pH 6.5 of Nimesulide. A higher percentage of drug was released with 1% SLS already reported [8]. In F-InR1, F-InR2 and F-InR3 the average drug release were nearly 10% reduced in comparison to immediate release formulations. Reason may be due to concentration of HPMC which showed the controlled release behaviour when used in the range of 5–15% [21–23]. Hydrophilic polymer HPMC K4M showed inverse relationship with release rate of Nimesulide due to the presence of carboxylic group and swelling nature [21, 24]. In case of slow release formulations higher concentration of HPMC K4M (15–35%) along with poor solubility of Nimesulide further decreased 10% release rate as compared to the intermediate release formulation. Fig 1 showed the percentage release of Nimesulide 100 mg in fasted gastric and intestinal fluids. Immediate release formulations were compared with innovator brand and results revealed that F-IR1 showed highest similarity ($f_2$) values i.e., 85.685, 90.177, 73.559 and 68.559% at pH 1.2, 4.5, 6.8 and 7.4 respectively. Similarity factor of F-InR1 and F-InR2 after comparing with F-InR3 which was selected as a reference formulation due to its excellent physicochemical properties were 55.462 and 59.672 in pH 1.2, 62.695 and 67.337 in pH 4.5, 57.521 and 74.900 in pH 6.8.
and 60.188 and 80.791 in pH 7.4. Similarly, difference factor of F-InR1 and F-InR2 after comparing with F-InR3 were 12.947 and 10.005 in pH 1.2, 6.331 and 5.167 in pH 4.5, 8.999 and 7.445 in pH 6.8, 13.042 and 7.531 in pH 7.4. Similarity ($f_2$) values of F-SR1 and F-SR3 with F-SR2 were 76.891 and 68.195 in pH 1.2, 72.748 and 55.309 in phosphate buffer pH 4.5, 74.900 and 58.828 in slightly acidic pH 6.8, 80.791 and 45.697 in slightly basic pH 7.4. Formulations F-IR1 from immediate release, F-InR2 from intermediate release and F-SR3 from the slow release were selected for further pharmacokinetic studies due to their excellent physicochemical and quality control attributes. Table 2 showed the correlation ($r^2$) values of different release formulations in different dissolution apparatuses and dissolution mediums.

### 3.2 In vivo pharmacokinetic studies

For the log-transformed values, the geometric mean ratio of $C_{\text{max}}$ calculated for reference and test was 0.995 µg/ml while geometric mean $C_{\text{max}}$ values were 6.087 ± 1.072 and 6.060 ± 1.073 µg/ml respectively. ANOVA results were found to be in significant range. P values of formulations and periods were insignificant whereas sequence and subject were found to be significant. The 90% confidence interval (CI) for $C_{\text{max}}$ (observed) were (99.17–100.3%) and for $C_{\text{max}}$ (calculated) were (99.3–99.7%). Similarly, values of geometric mean for $T_{\text{max}}$ (observed) for reference and test formulations were (1.3812 ± 1.1745 and 1.366 ± 1.1724 hr). In ANOVA the formulations, subject and sequence were found to be insignificant having geometric ratio of 0.989. The 90% confidence interval values for $T_{\text{max}}$ were 98.198 to 99.658%. Observed values of $AUC_{\text{ext}}$ were 100.95 to 104.91% and $AUC_{\text{total}}$ were 99.88 to 100.08%. For $AUC_{\text{ext}}$ and $AUC_{\text{total}}$ values for geometric mean values for both reference and test products were (0.110 ± 1.378 and 0.1141 ± 1.363) and $AUC_{\text{total}}$ were (23.6451 ± 1.02917 and 23.640 ± 1.028) respectively. For non-log transformed data, values of geometric mean for $C_{\text{max}}$ (observed) of reference and test products were (6.01084 ± 1.19275 and 5.99586 ± 1.19614), while $C_{\text{max}}$ (calculated) were (6.087 ± 1.072 and 6.060 ± 1.073). Values of $AUC_{\text{total}}$, $T_{\text{max}}$ and $AUC_{\text{last}}$ were found within the required limits. For non-log transformed data, the 90% confidence intervals (CI) values for $C_{\text{max}}$ (observed) (99.25–100.32%), $C_{\text{max}}$ (calculated) (99.33–99.69%), $T_{\text{max}}$ (98.188–99.834%) $AUC_{\text{total}}$ (99.88–100.08%). In this study values of $C_{\text{max}}$, AUC and $T_{\text{max}}$ of F-IR1 and

### Table 2. Correlation ($r^2$) values using Weibull model for Nimesulide 100 mg tablets at different conditions of dissolution apparatus and different composition of dissolution mediums.

| Formulation | pH 1.2 | pH 4.5 | pH 6.8 | pH 7.4 | FaSSGF | FeSSIF | FaSSIF |
|-------------|--------|--------|--------|--------|--------|--------|--------|
| **USP Dissolution apparatus I with 100rpm** | | | | | | | |
| Reference Product | 0.897 | 0.878 | 0.888 | 0.894 | 0.914 | 0.884 | 0.842 |
| F-IR1 | 0.889 | 0.889 | 0.887 | 0.902 | 0.812 | 0.784 | 0.812 |
| F-InR2 | 0.885 | 0.912 | 0.895 | 0.912 | 0.842 | 0.841 | 0.874 |
| F-SR3 | 0.998 | 0.912 | 0.999 | 0.999 | 0.884 | 0.910 | 0.841 |
| **USP Dissolution apparatus II with 100rpm** | | | | | | | |
| Reference Product | 0.909 | 0.934 | 0.933 | 0.995 | 0.874 | 0.898 | 0.826 |
| F-IR1 | 0.996 | 0.988 | 0.995 | 0.988 | 0.888 | 0.876 | 0.888 |
| F-InR2 | 0.900 | 0.922 | 0.911 | 0.914 | 0.733 | 0.940 | 0.915 |
| F-SR3 | 0.989 | 0.990 | 0.997 | 0.993 | 0.884 | 0.873 | 0.986 |
| **USP Dissolution apparatus II with 50rpm** | | | | | | | |
| Reference Product | 0.785 | 0.898 | 0.888 | 0.952 | 0.841 | 0.884 | 0.914 |
| F-IR1 | 0.841 | 0.875 | 0.899 | 0.914 | 0.774 | 0.821 | 0.845 |
| F-InR2 | 0.712 | 0.897 | 0.912 | 0.8965 | 0.812 | 0.814 | 0.842 |
| F-SR3 | 0.999 | 0.998 | 0.956 | 0.9458 | 0.741 | 0.941 | 0.924 |

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3.3 Development of IVIVC correlations

Multiple formulations of F-IR1, F-InR2 and F-SR3 were designed and developed using HPMC as a rate controlling polymer. Data indicated that the as the concentration of HPMC reduces the drug release rate. In this study, in vitro release assessment were conducted at several dissolution media i.e. pH 1.2, 4.5, 6.8, 7.4; FaSSGF, FaSSIF and FeSSIF using USP dissolution Apparatus I (at 100 rpm) and Apparatus II (at 50 and 100 rpm). Fig 2 showed the development of Level A IVIVC using the deconvolution method for Nimesulide 100 mg tablets. At pH 1.2 (50 rpm of USP II and 100 rpm of USP II and I), numeric values of % PE of $AUC_{inf}$ were found to be (6.788, 9.587 and 8.974) and $C_{max}$ (6.055, -2.414 and 1.531) respectively. Similarly, numeric values of %PE of $AUC_{inf}$ at pH 4.5 and pH 6.8 were found to be (9.156, 9.207 and 9.355) and (2.584, 2.978 and 2.472) and $C_{max}$ values were (-4.745, -3.276 and -5.526) and (2.584, 0.573 and 4.692) respectively. At pH 7.4 (50 rpm of USP II and 100 rpm of USP II and I), consecutive numeric values of $AUC_{inf}$ and $C_{max}$ were found to be (4.350, 7.173 and 7.094; 5.621, 3.055 and -2.471). Similarly, values of % PE of $AUC_{inf}$ were consecutively found to be (9.372, 9.542 and 9.150); (9.900, 9.167 and 8.147) and (9.886, 8.212 and 9.130) and $C_{max}$ (-7.228, 8.957 and -6.680); (3.695, 7.573 and 9.161) and (-29.059, -6.085 and -7.061) at FaSSGF, FaSSIF and FeSSIF (50 rpm of USP II and 100 rpm of USP II and I) as shown in Table 3. Eddington et al., in 1998 reported values of internal prediction error (%PE) of $C_{max}$ and $AUC_{last}$ of metoprolol tartrate (ER) tablets [25].

For $AUC_{inf}$ and $C_{max}$, the %PE value of each formulation was found to be < 15% while the average values were < 10%. The low values of prediction errors demonstrate that the correlation
model is projecting the in vivo response of each formulation. Thus indicated a valid Level A IVIVC according to the FDA guidance document [26]. External (%PE) was not required because individual values of internal (%PE) error of \(C_{\text{max}}\) and AUC last were not >15. As compared with other nonlinear models, the Weibull model was considered to be the best fit for in vitro-in vivo data. Estimation of regression values \(r^2\) is significantly important for point to point correlation. Correlation \(r^2\) values after comparison of in-vivo absorbed vs. in-vitro drug dissolved values at different dissolution media were successfully calculated. Linear correlation found at pH 6.8 and 7.4 i.e. \(r^2 = 0.999\). Jantratid et al in 2006 reported excellent value of correlation i.e. \(r^2 = 0.968\) for cimetidine tablets to develop IVIVC model [27]. The reported results are quite similar to the Tandt et al., in 1995 reported the IVIVC correlation at phosphate buffer pH 6 [28]. Results of the present study indicated that IVIVC correlation was obtained at pH 6.8 and pH 7.4 showed good correlation Also, internal prediction error (%PE) of AUC and \(C_{\text{max}}\) were calculated as presented in Table 3. In the previous literature, the fed state of gastric and intestinal medium also considered as the best for correlation[18, 19]. Leu et al in 2008 established the IVIVC model for hemibenzathine and demonstrated that best medium was found to be fed state to predict AUC and \(C_{\text{max}}\) showed <10% prediction error [29].

**Conclusion**

In this study, a Level A IVIVC was established which demonstrate the relationship between in vivo absorption data and in vitro release data for all Nimesulide products. Results of internal validation were found to be within the adequate limits indicating the prediction of correlation models.

**Supporting information**

S1 File. “Comparative Excel Data of Test and Reference Brand.xlsx”.
(XLSX)

S2 File. “In vivo data IVIVC.xlsx”.
(XLSX)

**Author Contributions**

Conceptualization: Muhammad Harris Shoaib.

Formal analysis: Muhammad Harris Shoaib.
Investigation: Rabia Ismail Yousuf.
Methodology: Muhammad Hanif.
Software: Rabia Ismail Yousuf.
Validation: Farya Zafar.
Visualization: Farya Zafar.

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