Quality Criteria Establishment for Dissolution of Ascorbic Acid from Sustained Release Pellets

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Abstract

With the advancement in the manufacturing industry of the food additive and supplements coupled with ever-increasing challenges in the health risk hazards, the establishment of strict control measures become essential to ensure product quality and safety. A small firm for manufacturing of food supplements and additives established a production line for a new product based on L-ascorbic acid (vitamin C) 500mg per hard gelatin capsule. Ascorbic Acid is provided as sustained release pellets. One of the important quality control tests that should be inspected was the dissolution profile after one, four and eight hours. Dissolution rate test was performed for 70 consecutive batches and results were interpreted. The best-fitting model that described the dissolution rate was Morgan-Mercer-Flodin (MMF) Model which is Sigmoidal four parameter mathematical curve. Establishment of Control Limits (CLs) was determined for each chronological test point using commercial statistical software package. Control limits standard deviations (sd) for one, four and eight hours were 28.1%±3.5%, 68.1%±4.1% and 89.4%±3.3%, respectively. On the other hand, the calculated lower and upper control limits (LCL, UCL) were (21.1%, 35.1%), (59.3%, 77.0%) and (81.6%, 97.2%), respectively. Interestingly, no Out-Of-Specification (OOS) results were detected for individual lots of product, yet UCLs exceeded Upper Specification Limits (USLs). This shift in the dissolution rate process is evident, especially in four hours point. Within/Overall Capability plots confirmed this finding with strong shift was detected to the right side that indicated a crucial need for investigational analysis. Accordingly, Statistical Process Control (SPC) provided provision for early warning of deviations in the manufacturing process cycle of the product before any true excursions may occur. The selection of product that has simple manufacturing processsing steps such as Vitamin C will aid in the spotting of the most probable causes of hidden deviations which may be related to consistency of raw materials quality and/or laboratory system related-deviations.

Keywords: Ascorbic acid; Hard gelatin capsule; Pellets; Dissolution rate; MMF model

Abbreviations: MMF: Morgan Mercer Flodin; CL: Control Limit; LCL: Lower Control Limit; UCL: Upper Control Limit; USL: Upper Specification Limit; SPC: Statistical Process Control; HGC: Hard Gelatin Capsule; API: Active Pharmaceutical Ingredient; QC: Quality Control; QA: Quality Assurance

Introduction

Many animals can synthesize vitamin C, which is known also as L-ascorbic acid, endogenously. However, the human body, in contrast to these organisms, has not the ability to synthesize this compound. Accordingly, it may be acquired from exogenous source as food supplement [1]. Vitamin C is involved in protein metabolism in addition to the synthesis of several biomolecules such as L-Carnitine, collagen and certain neurotransmitters [1,2]. Moreover, L-ascorbic acid acts as an antioxidant and has an important role to regenerate other antioxidants within body such as α-tocopherol, which is also called vitamin E [3,4]. In addition, L-ascorbic acid has a crucial function in the immune system [3] and enhance the absorption of non-heme iron [5]. Due to the physiological importance of vitamin C, deficiency in its daily intake causes disease condition known as “Scurvy”, with prominent signs include: lassitude, fatigue and fragile capillary system associated with widespread connective tissue weakness [6-9].

Due to the clinical and physiological importance of vitamin C, it has been incorporated in different forms by the manufacturers of the medicinal drug products [10]. During the last decades, extended release formulations became more common and popular because they offer several advantages over the classical dosage forms in terms of financial costs, toxic effects, dosing regimen and instability of blood plasma concentrations [11]. Efficiently low but therapeutically effective plasma concentrations of the active medicinal products can be achieved usually between 8 and 12 hours coupled with minimized risk of local irritation are among major advantages of the prolonged release drug forms [12-15].

However, the efficacy and quality of the medicinal product delivered to the final customer is largely dependent on the monitoring of certain inspection properties using statistical tools. Statistical Process Control (SPC) may be defined as the utilization of factual techniques to check and monitor current procedure to
guarantee that it works at its maximum capacity to create acceptable units. Under SPC, a procedure carries on typically to create however much accommodating item as could reasonably be expected with the minimum conceivable waste. While SPC has been connected most as often as possible to the controlling of manufacturing and production lines, it applies similarly well to any procedure with a quantifiable yield. Using SPC techniques will allow for triggering continuous enhancement projects [16].

Since the final consumable products affect human health directly, monitoring of manufactured batches should be strictly controlled in terms of quality and consistency. The present study aimed to apply SPC on one of the important inspection characteristics of L-ascorbic acid sustained release pellets in form of Hard Gelatin Capsule (HGC) viz. dissolution rate of Active Pharmaceutical Ingredient (API) at predefined intervals. A newly installed plant for manufacturing of food supplements and additives has launched this vitamin C product and performed long-term inspection about the current state of quality of the manufactured product and the level of improvement needed to ensure compliance and stability of the dosage form quality characteristics.

Materials and Methods

A recently established plant for food supplements and additives manufacturing has launched a new product as HGC containing sustained release vitamin C pellets. Detailed data of dissolution test of 70 successive batches of the product from Quality Control (QC) laboratory were collected and processed by the quality team of the firm - using commercial statistical software packages - to investigate the current state of the firm compliance to GxP and the degree of improvement required. This work is part of total quality characteristic monitoring project that guides the industry futuristic development. GxP refers to Good (G), broad set of quality references and guidelines (x) and Practice (P) (International Society for Pharmaceutical Engineering [17].

The Content of each Capsule includes 600mg of vitamin C pellets with a net content of ascorbic acid equivalent to 500mg. Other inactive components are beeswax, stearic acid and microcrystalline cellulose. The Dissolution test was performed for 70 batches based on an internal method developed by the manufacturer of the sustained release pellets. Dissolution medium was made of buffer pH 2.9, volume 900ml and in USP paddle apparatus with speed 100rpm.

Figure 1: Simplified manufacturing process diagram of ascorbic acid capsule.

Simplified manufacturing processing steps were illustrated graphically as shown in (Figure 1) Determination of the process limits and parameters: Centre line and control limits are calculated as detailed literally as detailed in Minitab [18]. Determination of best fitting model for curve was determined using Curve Expert Version 1.4. Correlation analysis, outlier’s detection and column statistics were executed using Graph Pad Prism Version 6.01 for Windows.

Result and Discussion

Results of the dissolution test for 70 batches of vitamin C HGC are shown in (Figure 2) with out-of-control batches are indicated with red points. Figure 3 shows concurrent comparative dissolution pattern of the pellets at different testing intervals with slightly decreasing trend over time. It should be noted that no outliers were detected using ROUT (at Q = 10.0%) which was recommended method by Graph Pad Prism. Data did not show any Out-Of-Specification (OOS) results.

Kinetics of dissolution of L-ascorbic acid can be expressed by the following model:

Morgan-Mercer-Flodin (MMF) Model (Sigmoidal four parameter mathematical curve):

$$y = \frac{(a \cdot b + c \cdot x^d)}{b + x^d}$$

Where: Coefficient Data for control limit (CL) calculation are the following: $a = 1.41E-011$, $b = 3.64E+000$, $c = 1.30E+002$ and

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d =9.99E-001. y =Fraction of vitamin C dissolved after x hours. Coefficient Data for lower specification limit (LSL) determination are: a =5.38E-002, b =1.15E+004, c =2.28E+005 and d =6.71E-001. On the other hand, upper specification limit (USL) coefficient data are a =-2.30E-009, b = 4.48E+000, c =1.92E+002 and d =7.62E-001. On the other side, LCL and UCL coefficient parameters are a =-1.85E-011 and a =6.40E-011, b =4.82E+000 and b =2.84E+000, c =1.23E+002 and c =1.35E+002 and d =1.09E+000 and d =9.62E-001, respectively. They are presented graphically in Figure 4.

Figure 2: Trending charts showing the general dissolution profile after one, four and eight hours of 70 batches of Ascorbic Acid pellets with mean, UCL and LCL indicated (red spots are out-of- control points).

Figure 3: Concurrent comparative study of dissolution profile of 70 batches of Ascorbic acid showing general decreasing trend line (solid black line).
Control limits±Standard Deviations (sd) for one, four and eight hours were 28.1%±3.5%, 68.1%±4.1% and 89.4%±3.3%, respectively. On the other hand, the calculated lower and Upper Control Limits (LCL, UCL) were (21.1%, 35.1%), (59.3%, 77.0%) and (81.6%, 97.2%), respectively. These findings are illustrated graphically in Figure 4.

Table 1: Correlation between three successive dissolution rate testing points of L-ascorbic acid 500mg capsule: after one hour, four hours and eight hours.

| Spearman correlation Matrix | Correlation Coefficient of Dissolution Rate |
|-----------------------------|---------------------------------------------|
|                             | After One Hour     | After Four Hours | After Eight Hours |
| P value of Dissolution Rate | After One Hour     | 0.59             | 0.39             |
| After Four Hours            | 8.71E-08           |                 |
| After Eight Hours           | 1.00E-03           | 7.73E-13         |

Significant correlation between dissolution rates in the following descending order could be concluded from Table 1: four-eight hours > one-four hours > one-eight hours. Thus, dissolution profile of adjacent points are related more than distant ones.
Figure 5 demonstrates overall/within capability plot for the three sections of the dissolution test. Figure 5B shows strong shift of the process to the right side i.e. towards USL. The greater spreading of the overall over within dissolution process indicates unstable and unsatisfactory performance over long run assessment study.

Statistical quality control is a strategy by which organizations collect and break down gathered information on the fluctuations which happen within processing activity time with a specific end goal to figure out whether improvements are required [19]. Accordingly, interpretation of results in Figure 2 was subjected to SPC analysis to monitor its quality and improvement required rather than reliance on reporting individual observation results only which did not show either Out-Of-Specification (OOS) nor outlier values from the test.

MMF model of dissolution in the current study, that was shown in the curve of Figure 4, is different from the pattern shown by other researchers [15, 20]. This is an area for extended research to elucidate the variation in the dissolution due to pellets engineering.

Overall capability must be close enough to short-term one to ensure process stability and predictability and at the same time drifting to either limiting boundaries of the specification should be avoided. Moreover, CL should be contained within SL of the process [21, 22]. So, the process did not show stability over long run with CL exceeded SL indicating that improvement is required. In addition, strong shift toward higher rate was evident at four hours point of the dissolution test. The pellets have been obtained from the same source during the manufacturing of 70 batches after QA investigation of the facility. Nevertheless, some investigators had demonstrated that the release of ascorbic acid is different between individual pellets which in turn are affected by unique microscopic surface defects on each unit [23]. Thus, the manufacturing of controlled release materials should be under the condition of statistical control to produce a consistent outcome in the dissolution study.

Little influence on the product quality could be impacted by production because the whole manufacturing steps are automated as demonstrated in Figure 1. Possible influential factor from automatic filling is the average filling weight, which was found to show low correlation with the dissolution pattern suggesting that it has low (if any) effect on the true stability of the dissolution rate. Based on these findings, further investigation is required to determine the root cause of such state of instability which may be due to the supplier quality issue and consistency of the pellets and/or laboratory analysis problems which may be related to instruments, methods and/or analysts training.

Conclusion

SPC provides an important tool to assess the quality of the inspection characteristics and provisionally predicts the compliance of the underneath involved processes to GxP rules. In the present study, the automated manufacturing stages coupled with the absence of interfering processing steps that influence the quality parameters of the final product limit the questionable sources of out-of-control and instability of the investigated process. In the current case, the most probable cause is quality control good laboratory practice (GLP) issue with the possibility of unstable dissolution pattern that may be originated from the manufacturer of the sustained release pellets.

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