Robust Batch Process Scheduling in Pharmaceutical Industries: A Case Study

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Abstract—Batch production plants provide a wide range of scheduling problems. In pharmaceutical industries a batch process is usually described by a recipe, consisting of an ordering of tasks to produce the desired product. In this research work we focused on pharmaceutical production processes requiring the culture of a microorganism population (i.e. bacteria, yeasts or antibiotics). Several sources of uncertainty may influence the yield of the culture processes, including (i) low performance and quality of the cultured microorganism population or (ii) microbial contamination. For these reasons, robustness is a valuable property for the considered application context. In particular, a robust schedule will not collapse immediately when a cell of microorganisms has to be thrown away due to a microbial contamination. Indeed, a robust schedule should change locally in small proportions and the overall performance measure (i.e. makespan, lateness) should change a little if at all.

In this research work we formulated a constraint programming optimization (COP) model for the robust planning of antibiotics production. We developed a discrete-time model with a multi-criteria objective, ordering the different criteria and performing a lexicographic optimization. A feasible solution of the proposed COP model is a schedule of a given set of tasks onto available resources. The schedule has to satisfy tasks precedence constraints, resource capacity constraints and time constraints. In particular time constraints model tasks due dates and resource availability time windows constraints. To improve the schedule robustness, we modeled the concept of \((a, b)\) super-solutions, where \(a, b\) are input parameters of the COP model. An \((a, b)\) super-solution is one in which if \(a\) variables (i.e. the completion times of a culture tasks) lose their values (i.e. cultures are contaminated), the solution can be repaired by assigning these variables values with new values (i.e. the completion times of a backup culture tasks) and at most \(b\) other variables (i.e. delaying the completion of at most \(b\) other tasks). The efficiency and applicability of the proposed model is demonstrated by solving instances taken from a real-life pharmaceutical company. Computational results showed that the determined super-solutions are near-optimal.

Keywords—Constraint programming, super-solutions, robust scheduling, batch process, pharmaceutical industries.

I. INTRODUCTION

In pharmaceutical production industry, batch production typically requires the culture of a microorganism population (i.e. bacteria, yeasts or antibiotics). In this application context, the main sources of uncertainty are low performance and quality of the cultured microorganism population as well as microbial contamination, during preparation and fermentation phases. For these reasons, robustness is a key aspect for antibiotic production planning. When a cell of microorganisms has to be thrown away, a robust schedule remains feasible or should change locally in small proportions and the overall performance measure (i.e. makespan, lateness) should change a little if at all.

A classification of problems in terms of plant configurations for pharmaceutical industry is given by [1]. Due to the different plant configurations and process specifications, a wide variety of optimization models, mainly Mixed Integer Linear Programming (MILP) models, have been proposed in the batch process engineering literature. In [2] Maravelias and Grossmann proposed a general hybrid MILP/CP iterative algorithm for the scheduling of multipurpose plants exploiting the strong points of Mathematical and Constraint Programming. Graph based methods usually can be used in order to fit more to the combinatorial nature of scheduling problems. In contrast to the infeasibility problems which may arise with MILP based models, graph based methods never provide infeasible solutions [3]. In this research work we formulated a constraint programming optimization (COP) model for the robust planning of antibiotics production. We developed a discrete-time model with a multi-criteria objective, ordering the different criteria and performing a lexicographic optimization.

The paper is organized as follows. In Section 2 we describe the antibiotics production process in terms of phases and the required resources, taking into account of the priority among activities. In Section 3, we define the problem statements and we give a mathematical formulation for the problem.

II. ANTIBIOTICS PRODUCTION PROCESS

Although most antibiotics are available in nature, such microorganisms are not normally produced in the quantities necessary for large-scale demand. For this reason, each pharmaceutical industry has to design and implement a specific process, typically referred to as fermentation process [4], for each antibiotic. This process (in Fig. 1) involves isolating a desired microorganism, fueling growth of the culture and refining and isolating the final antibiotic product. It is important that sterile conditions be maintained throughout the manufacturing process, because contamination by foreign microbes will ruin the fermentation.

A. Fermentation Process Description

The antibiotic is the output of a production process consisting of four main activities described in the following. In order to better understand the relations between the activities and the resources, we report in Fig. 2 an IDEF0 model. The fermentation process takes place using a culture medium.
and a charge of living matter. The culture medium and the charge of living matter are firstly prepared by a mixer and a pre-fermentor, respectively (preparation). The prepared culture medium is firstly transferred to a tank for dilution, then passed through a sterilizer and finally loaded into a fermentor using a load collector (culture medium inoculum). Once the culture medium is cooled (cooling), the pre-fermented charge is transferred from the pre-fermentor to a fermentor through a transfer collector (living charge inoculum). Then, the growth of the obtained culture is monitored and fueled (fermentation) and, after a given amount of time, the antibiotic is isolated (harvest).

Fig. 1. Fermentation process

Fig. 2. IDEF0 process model

III. PROBLEM STATEMENTS AND MATHEMATICAL MODELING

We modeled antibiotic production planning as a project scheduling problem as follows. We supposed that the planning horizon is partitioned into $N$ time slots. We denoted with $F$ the set of antibiotic types (i.e. project types), and $K$ the set of resource types (i.e. fermentor, sterilizer, etc.). For each antibiotic type $f \in F$, we defined a work breakdown structure, modeled as a directed graph $G_f = (T_f, P_f)$ where $T_f$ and $P_f$ represent, respectively, the set of tasks and the precedence constraints. For each task $t \in T_f$, we defined a fixed duration $s_t$ and a set of requirements $\Pi_t \subseteq K$, such that if $k \in \Pi_t$, then the resource type $k$ is required to execute task $t$. Each task $t$ has a sets of time windows $W^t_s (W^t_e)$, representing the feasible values for its start time (end time). A boolean parameter $m_t$ states if the execution of task $t$ is mandatory or not.

The set $R$ represents the available resources and each $r \in R$ is associated with a resource type $k_r \in K$. For each resource $r \in R$, it is defined a set $B_r$ of forbidden time windows, during which resource $r$ is unavailable (i.e. maintainance). We supposed a non-preemptive operating mode and a unit processing capacity.

A set of products orders $O$ represents the antibiotics demand, in particular each $o \in O$ refers to only one antibiotic type $f_o \in F$. Finally for each order $o \in O$, it is defined an earliest (latest) $d_{o}^{\min}$ ($d_{o}^{\max}$) ending time for the harvest task.

In the following we refer to $d_{o}^{\max}$ as the due date of order $o \in O$.

A. A Constraint Programming Model

Let $alt_{o,tr}$ be an interval variable representing the starting and ending times of task $t$, if assigned to resource $r \in R$, with $o \in O$, $t \in T_{f_o}$ and $k_r \in \Pi_o$. The proposed constraint programming model aims to determine the optimal values of $alt$ variables, minimizing the total lateness with respect to order due dates. In order to model lateness and precedence constraints, we denote with:

- $\text{start}(\text{var})$ the starting time for a given interval variable $\text{var}$;
- $\text{end}(\text{var})$ the ending time for a given interval variable $\text{var}$;
- $\text{presence}(\text{var})$ a boolean value equal to true if the interval variable $\text{var}$ is scheduled, false otherwise.

Let $\text{task}_{o,j}$ interval variable, representing the start and end processing times of task $t \in T_{f_o}$, $o \in O$. Let $\delta_o$ denote the lateness of order $o$, that is

$$\delta_o = \max \{0, \text{end}(\text{task}_{o,H_o}) - d_{o}^{\max}\},$$

where $H_o$ is the harvest task for the order $o \in O$.

Given an antibiotic type $f \in F$, precedence constraints have been partitioned into three types i.e. $P_{f_i} \cup P_{f^2} \cup P_{f^3} = P_f$. Each type of precedence constraints corresponds to one of the following set of linear constraints.

- $\text{StartAtEnd constraints}$: for each $(i,j) \in P_{f_i}$, task $j$ has to be started exactly $\tau_{ij}$ time units after the ending of task $i$, that is
  $$\text{end}(i) + \tau_{ij} = \text{start}(j).$$

- $\text{StartAfterEnd constraints}$: for each $(i,j) \in P_{f^2}$, the minimum (the maximum) delay between the end of task $i$ and the start of task $j$ is $\tau_{ij}^{\min}$ ($\tau_{ij}^{\max}$):
  $$\text{end}(i) + \tau_{ij}^{\min} \leq \text{start}(j) \leq \text{end}(i) + \tau_{ij}^{\max}.$$

- $\text{StartAfterStart constraints}$: for each $(i,j) \in P_{f^3}$, the minimum (the maximum) delay between the start of task $i$ and the start of task $j$ is $\tau_{ij}^{\min}$ ($\tau_{ij}^{\max}$):
  $$\text{start}(i) + \tau_{ij}^{\min} \leq \text{start}(j) \leq \text{start}(i) + \tau_{ij}^{\max}.$$

Given a resource $r \in R$, we modeled changeover times as follows. Firstly, we defined a decision variable $\sigma_{r,n}$ denoting...
the state of resource \( r \in R \) during time slot \( n \in N \). In particular, a resource \( r \in R \) can be either ready to process an antibiotic type, i.e. \( \sigma_{rm} = f \in F \), or not available due to setup operations, i.e. \( \sigma_{rm} = \text{nil} \). Let \( F = F \cup \{\text{nil}\} \) be the domain of \( \sigma_{rm} \), with \( r \in R \) and \( n \in N \). The setup cost \( M_{r \ell} \) represents the setup time required if a state transition occurs from \( \ell \in F \) to \( \ell' \in F \) on resource \( r \in R \).

In order to generate fault tolerant solutions and ensure robustness, we follow the reformulation approach suggested in [5] to build a super-solution.

A solution to a Constraint Satisfaction Problem is \((a, b)\) super-solution if the loss of the values of at most a variables can be repaired by assigning other values to these variables, and modifying the assignment of at most b other variables.

In this work we build super-solutions via reformulation with the duplication of some variables. The duplicate variables have the same domain as the original variables, and are linked by the same constraints. The assignment to the original variables is a super-solution, where the repair for each variable is given by its duplicate (see Fig. 3).

First objective function (1) is the total lateness, whilst second objective function (2) is the robustness. Constraints (3), (4) and (5) are the precedence constraints described in the previous section. For each order \( o \in O \), constraint (6) states that the end of the harvest task \( H_o \) has to occur in \([a_o^m, a_o^M + \delta_o]\). Constraint (7) models the relationship between the \( alt \) and the \( task \) variables. Constraint (8) represents the unit processing capacity for each resource \( r \in R \). Constraints (9) and (10) model time windows \( W_r^l \) and \( W_r^u \). Constraint (11) defines the domain of the \( task \) variables, while constraint (12) states their duration. Constraint (13) imposes the schedule of mandatory operations. For each resource \( r \in R \), constraint (14) models the forbidden time windows \( B_r \). Constraints (15) and (16) model state transitions applying the changeover times. We consider by convention that the changeover time from or to the nil status is equal to zero. Constraints (17) and (18) state respectively the domain and the initial value of states variables. Constraint (19) states an upper bound for the lateness.

IV. COMPUTATIONAL RESULTS

The efficiency and applicability of the proposed model was demonstrated by solving instances taken from a real-life pharmaceutical company. In Fig. 4 and 5 we reported the Gantt charts corresponding to the output of one instance. We used one different color for each \( f \in F \). On the left, the labels refer to resources \( r \in R \). In particular, labels \( F_r \) refer to fermentors that are the main bottlenecks (see Fig. 4).

When our research team began to deal with this decision problem, the production planning was determined by applying an earliest due date heuristic (EDD). The production manager was wondering if their monthly throughput could be increased.
The computational results pointed out that throughput can be increased only by adding one more fermentor. It is worth noting that adding one fermentor is a strategic decision since it requires huge investments (about one million dollars).

As far as robustness is concerned, Fig. 5 reports the Gantt chart obtained when we optimized in lexicographical order lateness and robustness: seven backup charges were added with respect to solution of Fig. 4. In general, our computational results showed that the proposed model was equivalent in terms of robustness, but provided a gain on lateness of 20% on average with respect to EDD heuristic. Finally, the EDD heuristic has a granularity of a day, whilst our model provides hourly details on the operational activities.

V. CONCLUSIONS

This work presents a constraint programming framework for the antibiotics production planning problem. We proposed a model for minimizing in lexicographical order total lateness and robustness, taking into account precedence, time and resource capacity constraints. Instances from a real-life pharmaceutical company showed a gain on lateness of 20% on average. The adoption of an automated tool for production scheduling is a way to improve productivity and to promote integration among strategic and operational decision making layers. This kind of vertical integration is a key step of Industry 4.0 [6]. In particular, this research work showed that robust constraint programming is an effective and efficient tool for planning industrial production of microorganism cultures.

REFERENCES