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AN INTERACTIVE TOOL FOR THE OPTIMISATION OF FREEZE- DRYING CYCLES BASED ON QUALITY CRITERIA

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Abstract

Among existing dehydration methods, freeze-drying has unique benefits for the stabilisation and preservation of biological activity of pharmaceutical products, but remains an expensive and time-consuming process. A user-friendly software tool was developed, allowing for interactive selection of process operating condition profiles in order to maximise process productivity while insuring product quality preservation. The software is based on a dynamic, one-dimensional heat and mass transfer model, which can accurately represent both the primary and secondary drying stages and the gradual transition between them. The model was validated in a wide range of operating conditions: -25 to +25°C shelf temperature and 10 to 34 Pa total pressure. By comparing a reference sucrose solution with a formulated pharmaceutical product containing polyvinylpyrrolidone (PVP), it is shown that controlling product properties such as glass transition temperature and sorption isotherm can reduce the minimum achievable cycle duration by 12 h (33%).

Keywords: lyophilization, dynamic modeling, optimization, glass transition temperature.

INTRODUCTION

In pharmaceutical industry, freeze-drying is widely used to preserve proteins and polypeptides, which are physically and/or chemically unstable in aqueous solutions (1). The improved stability of freeze-dried proteins provides many benefits such as storage stability at ambient temperature, extended shelf life, convenient handling

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and transport. However, freeze-drying is known to be a time-consuming and expensive process. An over conservative freeze-drying cycle results in long process time, added production costs, unnecessary consumption of drying capacity, and may compromise protein stability. Various process variables affect the efficiency of the freeze-drying process. The product temperature is a determinant factor for both productivity and product quality (2-4). The sublimation is generally faster at higher temperatures, but drying at excessive temperatures results in a loss of the pore structure obtained by freezing, which is defined as the collapse phenomenon by Pikal and Shah (5) and therefore in rejection of the batch. For amorphous formulations, the collapse is usually associated with the glass transition temperature of the maximally freeze-concentrated phase. The product temperature results from the shelf temperature and the chamber pressure applied and is not directly controlled during freeze-drying process. In practice, the appropriate shelf temperature and chamber pressure conditions are frequently established empirically in a “trial-and-error” experimental approach and many manufacturing processes are far from optimal (6). Theoretical modelling can provide a better understanding of the impact of process and formulation variation on cycle time and product temperature history and, thereby, facilitate process development and on-line monitoring (7).

This study presents a software tool for interactive selection of the process operating conditions: shelf temperature profile, chamber pressure profile, condenser temperature, etc. While changing the operating conditions, the user monitors process parameters relevant for product quality and productivity: product temperature and glass transition temperature at critical points in order to assess product stability, as well as the residual moisture content which determines the cycle duration. Unlike fully automated optimisation (8, 9), the user retains complete control over the selected operating conditions, which can be advantageous for gaining insight into the limiting factors and for taking into account semi-empirical, experience-based considerations, difficult to encode in a formal optimisation process. The software is based on a simple yet accurate one-dimensional dynamic model of heat and mass transfer with associated product quality indicators. The development of this model was motivated by the necessity of very quick simulation compared to many existing complex multi-dimensional models (10-13). Short simulation time, less than 0.2 s on a standard PC (AMD Athlon[®] processor at 2 GHz), allows comfortable interaction with the user. Detailed output information is also available for posterior analysis under Matlab[®] and Excel[®], such as temperature, vapour pressure, residual moisture profiles in relevant locations in the product layer and in the chamber.

MATERIALS AND METHODS

Experimental

Product formulations

Two product formulations were considered in this study. The first formulation (coded S) was a 5% sucrose and 10 mM Tris-HCl solution taken as reference. The second formulation (coded PS) contained 4% polyvinylpyrrolidone (PVP), 1% sucrose and 10 mM of Tris-HCl and was selected for its efficient protein stabilisation properties (14). Sucrose was purchased from Prolabo (Paris, France) and polyvinylpyrrolidone (with a reported average molecular weight of 25,000 Da) from Merck (Darmstadt, Germany).

Freeze-drying protocols

One millilitre of each formulation was placed in 4 mL glass vials. The vials (a total of 213 vials per cycle) were placed on an aluminium tray and loaded onto the shelf of a SMH 15 freeze-drier (Usifroid, Maurepas, France). The cycle consisted of freezing at -45°C (shelf cooling rate at $0.6^{\circ}\text{C}/\text{min}$ and holding at -45°C for 2h), followed by primary drying under various conditions of shelf temperature (-5°C , -15°C , -5°C , 5°C , 15°C and 25°C) and chamber pressure (10 Pa, 18 Pa, 26 Pa and 34 Pa) and, finally, a secondary drying of 6 hours at 25°C and 10 Pa. Depending on the experiment, the heating rates of the shelf temperature were of $0.1^{\circ}\text{C}/\text{min}$, $0.25^{\circ}\text{C}/\text{min}$ or $1^{\circ}\text{C}/\text{min}$.

Product temperature was measured by two thermocouples placed at the bottom of two different vials. A moisture sensor (Panametrics Ltd, Shannon, Ireland) was used to monitor the partial vapour pressure in the chamber.

Lyophilized samples were equilibrated at 25°C under the following saturated salt solutions of constant water activities (a_w values are shown in parenthesis): CH_3COOK (0.23), $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$ (0.33), K_2CO_3 (0.44), $\text{Mg}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ (0.53), NaCl (0.75) and KCl (0.84).

Differential scanning calorimetry

DSC was used to determine the glass transition temperature of liquid samples before freeze drying, and samples that were freeze dried and equilibrated to different relative humidities. DSC measurements were performed using a power compensation differential scanning calorimeter (DSC) (Pyris 1 model; Perkin Elmer LLC, Norwalk, CT, USA), equipped with a liquid nitrogen accessory (CryoFill, Perkin Elmer). Between 5 and 20 mg of each

sample were placed in 50 μL Perkin-Elmer DSC aluminium sealed pans, and an empty pan was used as a reference. Cooling and heating rates of $10^\circ\text{C}/\text{min}$ were used. The glass transition temperatures were reported as the midpoint temperature of the heat capacity step associated with the glass transition, in accordance with the ASTM Standard Method E 1356-91.

Liquid samples were cooled to -120°C to ensure temperature stability and sample equilibration, and scanned for the first time to 25°C . This conventional thermal cycle was replicated once to verify the absence of time dependent changes in thermal behaviour. Dried samples were cooled to -40°C and then heated to 200°C .

Water content determination

Water content of the freeze-dried product and the samples equilibrated under different relative humidity conditions was measured by the Karl Fisher titration method using a Metrohm KF 756 apparatus. At least 20 mg of powder were mixed with 2 mL of dry methanol and titrated with Riedel-deHaen reagent (Seelze, Germany).

Water activity measurement

Water activity of samples from sorption isotherms at 25°C was measured by an aw meter FA-st/1 (GBX Scientific Instruments, Romans sur Isère, France). Measurements were made with dynamic method and each result was the average value of 10 measurements once the stability had been reached.

Dynamic model of the freeze-drying process

Assumptions

A conventional one-dimensional freeze-drying model of heat and mass transfer was developed, based on the following assumptions:

- A1. The product and the dryer state is described by relevant state variables in six key points shown in Figure 1: shelf, product bottom, sublimation front, product top, freeze-drying chamber and condenser. The heat and mass transfers are assumed to take place between these points.
- A2. Slow dynamics such as sublimation front movement and residual water desorption are described explicitly by differential equations. Relatively fast dynamics (compared to the typical duration of a freeze-drying cycle) such as heat transfer in the frozen and in the dry product layers, as well as the mass transfer in gaseous phase, are assumed to be in quasi-steady state and are described by algebraic equations. This assumption is supported by detailed calculation of the relevant time constants, reported in Appendix 1.
- A3. The residual water desorption flux was neglected in the heat and mass balances. A complete model, including the desorption heat and desorption vapour fluxes was also built and tested, and it was found that the contribution of the desorption was actually negligible, confirming earlier findings (15).

- A4. Ice gradually disappears in various product regions. The relative importance of the primary and secondary drying mechanisms is given by the ratio of the remaining sublimation area to the total product cross area.

Control variables

The shelf, chamber and condenser temperatures, as well as the total chamber pressure are assumed to be imposed by the freeze-drying protocol and known at any moment. They are allowed to vary in time, however.

State variables

The product state at any given moment is described by the sublimation front position, the temperature and the residual water concentration at the product bottom, front and top.

During the primary drying stage, the front temperature is determined from the heat balance condition at the sublimation front, taking into account assumptions A2 and A3:

$$\begin{aligned} & H^{Shelf\ Front}(Z^{Front}, P_t^{Chamb}) \cdot (\Theta^{Shelf} - \Theta^{Front}) + H^{Front\ Chamb}(Z^{Front}, P_w^{Sat}(\Theta^{Front})) \cdot (\Theta^{Chamb} - \Theta^{Front}) \\ & = L_{subl} K^{Front\ Cond}(Z^{Front}) \cdot (P_w^{Sat}(\Theta^{Front}) - P_w^{Sat}(\Theta^{Cond})) \end{aligned} \quad (1)$$

Eq. (1) expresses the fact that, in the assumed quasi-steady state situation, the net heat flux towards the sublimation front (from the product shelf and from the chamber) mainly serves for ice sublimation. In this equation, the heat transfer coefficients $H^{Shelf\ Front}$ and $H^{Front\ Chamb}$, as well as the mass transfer coefficient $K^{Front\ Cond}$ are calculated using resistance in series formulae (Figure 1). The heat transfer between the shelf and the product bottom includes contact, radiation and gas conduction terms. The heat transfer resistance through the glass vial bottom turned out to be negligible. The gas conduction term depends on the total pressure in the freeze-drying chamber (2). The heat transfer between the product bottom and the sublimation front takes place by conduction through the frozen layer, and hence depends on its thickness (front position). The heat transfer between the sublimation front and the chamber takes into account transfer through the dry layer to the product top (by gas conduction, vial walls and radiation) and from the product top to the chamber, mainly by radiation. The mass transfer resistance includes flow resistance through the porous dry layer (depending on its thickness), from product top to the freeze-drying chamber (specifically to the location of the partial vapour pressure sensor) and from the chamber to the condenser. These last two transfer coefficients depend on the freeze-dryer design. For sake of readability and completeness, detailed expressions for all heat and mass transfer coefficients are given in Appendix 2 in a tabular form.

Eq. 1 encodes the unavoidable interdependence between the heat and mass fluxes. The heat flux depends on the sublimation front temperature and determines the mass flux. The mass flux creates a vapour pressure increase in the dry layer, which in turn determines the sublimation front temperature, because of the local (at the sublimation

front) equilibrium condition between ice and vapour. In practice, solving Eq. 1 gives the sublimation front temperature.

The front position is determined from the sublimation mass flux by considering the volume liberated by the sublimated ice:

$$\frac{dZ^{Front}}{dt} = \frac{H^{FrontCond}}{D^{Ice} A} (P_w^{Sat}(\Theta^{Cond}) - P_w^{Sat}(\Theta^{Front})) \quad (2)$$

During the secondary drying stage, the front position is fixed to zero and the front temperature is formally taken equal to the product bottom temperature.

During both drying stages the residual water concentration at the top of the dry layer is calculated using a first-order desorption kinetic:

$$\frac{dC^{Top}}{dt} = \frac{1}{\tau_{Des}} \left[C_{Equ} \left(\frac{P_w^{Top}}{P_w^{Sat}(\Theta^{Top})} \right) - C^{Top} \right] \quad (3)$$

The equilibrium water content C_{Equ} is given by the sorption isotherm, as described in the following section. It depends on the ratio between the actual vapour pressure and the saturation pressure. At equilibrium, the ratio of these pressures would represent the water activity of the product. The actual vapour pressure is determined from the mass flux and the resistance in series model (Figure 1).

Similar equations were written for the product top and for the sublimation front. During the primary drying, at the product bottom and at the sublimation front, the vapour is in equilibrium with the ice according to assumption A2 and the pressure ratio is equal to one. During the secondary drying stage, the front position is formally identical to the product bottom and the partial vapour pressure is in equilibrium with the chamber and the condenser. This later condition results from assumptions A2 and A3: the desorption mass flux being negligible, there is no local pressure increase in the dry layer (16).

Finally, the model consists of a system of four differential equations (for the front position and the residual moisture contents at the product bottom, front and top) and one algebraic equation for the front temperature. The five coupled equations are solved simultaneously using the Matlab[®] numeric computation software (Natick, MA, USA). A detailed description of the model is given in the Appendix 2.

Output variables

The model allows a relatively straightforward calculation of various quantities of practical interest, in any relevant location among those shown in Figure 1, such as: product temperatures, glass transition temperatures

important for the product stability during drying, partial vapour and neutral gas pressures, average residual moisture content important for assessing the end of the drying cycle, evolution of the product mass in a vial, etc.

Gradual transition between the primary and the secondary drying stages

In practice, the freeze-drying conditions are not perfectly homogeneous (17). Lack of homogeneity appears both among vials, e.g. due to position on the shelf, wall chamber radiation, unequal vial filling etc. as well as inside each vial, e.g. due to heat conduction through vial walls, lack of planarity of the vial bottom, etc. At the scale of the freeze-dryer, the transition between the primary (I) and the secondary (II) drying stages appears gradual. According to assumption A4, this gradual “transition function” was defined as the ratio of the remaining sublimation area to the total product cross area:

$$\frac{A_{(I)}(t)}{A^{Tot}} = f(Z^{Front}(t)) \quad (4)$$

During the transition between stages, both models for the primary and the secondary drying are run in parallel, each with its own product cross section area:

$$A_{(I)} = f(Z^{Front})A^{Tot} \quad A_{(II)} = (1 - f(Z^{Front}))A^{Tot} \quad (5)$$

The reported global values of the relevant variables are averages, weighted according to the transition function:

$$\Theta^{Front} = f(Z^{Front})\Theta_{(I)}^{Front} + (1 - f(Z^{Front}))\cdot\Theta_{(II)}^{Front} \quad (6)$$

$$\frac{dZ^{Front}}{dt} = f(Z^{Front})\frac{dZ_{(I)}^{Front}}{dt} + (1 - f(Z^{Front}))\cdot 0 \quad (7)$$

$$\frac{dC^{Top}}{dt} = f(Z^{Front})\frac{dC_{(I)}^{Top}}{dt} + (1 - f(Z^{Front}))\frac{dC_{(II)}^{Top}}{dt} \quad (8)$$

Similar weightings are used for the other state and output variables.

The transition function was defined empirically as:

$$f(Z^{Front}) = \begin{cases} 1 & \text{if } Z^{Front} \geq Z^{Trans} \\ \sin\left(\frac{\pi}{2} \frac{Z^{Front}}{Z^{Trans}}\right) & \text{if } Z^{Front} < Z^{Trans} \end{cases} \quad (9)$$

Eq. 9 says that the primary drying is the only phenomenon as long as the sublimation front position Z^{Front} is above some critical transition value Z^{Trans} . Below this value, the relative importance of secondary drying increases until the sublimation front reaches the vial bottom ($Z^{Front} = 0$), when it becomes the only phenomenon.

The value of Z^{Trans} is a measure of the smoothness of the transition. The larger Z^{Trans} , the longer the transition between the primary and the secondary drying.

Models of the product properties relevant for cycle optimisation

Glass transition temperature

An important condition for product stability in the frozen layer during freeze-drying is absence of collapse (5). The product temperature must remain below the collapse temperature in any point during the whole freeze-drying cycle. The collapse temperature is usually about 2°C higher than the glass transition temperature of the freeze-concentrated phase Θ_{Glass}^{Frozen} for formulations that remains amorphous during freezing (5). The stability of the product in the dry layer is insured if the product temperature is maintained below the glass transition temperature (18), which strongly depends on the residual moisture content. The condition on the glass transition temperature was thus retained as the main product stability criterion for both the frozen and the dry layers. The evolution of the glass transition temperature was modelled by the classical Gordon-Taylor equation, slightly modified to take into account the frozen layer:

$$\Theta_{Glass}(C) = \max \left\{ \Theta_{Glass}^{Frozen}, \frac{(1-C)\Theta_{Glass}^{Dry} + q_{Gordon} C \Theta_{Glass}^{Ice}}{(1-C) + q_{Gordon} C} \right\} \quad (10)$$

where C is the moisture content, Θ_{Glass}^{Dry} and Θ_{Glass}^{Ice} are the glass transition temperatures of the perfectly dry product and of the ice respectively and q_{Gordon} is the Gordon-Taylor shape coefficient. For each product formulation, the Gordon-Taylor coefficient was determined by fitting the considered equation to experimental glass transition data of the samples equilibrated under various relative humidity conditions (Figure 2).

Sorption isotherm

A quality requirement on the final freeze-dried product is to reach a pre-specified residual moisture content, both under- and over-drying being damageable. Moreover, the glass transition temperature in the dry layer strongly depends on the moisture content. Both these conditions require the modelling of the water desorption process in the dry layer, which in turn requires the knowledge of the sorption isotherm and of the desorption kinetic.

None of the commonly used models (Freundlich, Langmuir, BET, GAB) was found to fit experimental sorption isotherm data adequately for the considered formulations. The fit was particularly inappropriate in the region of small water activities (less than 0.2) leading to clearly wrong predictions of the final product moisture content. A piecewise linear model was found to give satisfactory results:

$$C_{Equ}(a_w) = \min \left\{ C_{Max}, \max \left\{ C_{Min}, q_{Sorp1}(a_w - q_{Sorp2}) \right\} \right\} \quad (11)$$

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Here C_{Equ} is the equilibrium moisture content, a_w the water activity, and the other parameters are constants depending on the product formulation. The maximum adsorbed moisture content C_{Max} was determined from the glass transition temperature of the frozen product (Figure 2). The fit of this model to the experimental data at 25°C is shown in Figure 3.

Desorption kinetic

The calculation of the residual moisture content evolution in time requires the knowledge of the desorption time constant in addition to the equilibrium moisture content (sorption isotherm). The following exponential desorption kinetic model was used, reflecting a first-order rate desorption mechanism (15):

$$C(t) = C_{Equ} + (C_{ini} - C_{Equ})e^{-\frac{t}{\tau_{Des}}} \quad (12)$$

Here τ_{Des} is the desorption time constant. The fit of this model to the experimental data is shown in Figure 4.

RESULTS AND DISCUSSION

Model validation

The dynamic freeze-drying model was validated based the on-line measurements of product temperature and partial vapour pressure in the chamber. Additionally, the final average moisture content of the product was checked against the simulated value. The product-specific properties are listed in Table 1. The characteristics of the freeze-drying apparatus and other product-independent parameters are given in Table 2. The results of the model validation are shown in Figure 5 for the two extreme couples of operating conditions considered in the experimental design: shelf temperature of -25°C with total chamber pressure of 10 Pa, and shelf temperature of +25°C with total chamber pressure of 34 Pa during primary drying. The agreement between measurements and model predictions is quite satisfactory, taking into account the accuracy of the measurements and the repeatability of the experiments: $\pm 1^\circ\text{C}$ for product temperature, ± 2 Pa for vapour pressure and ± 0.01 kg/kg for the final moisture content.

Freeze-drying cycle optimisation

After the validation step, the model was used for freeze-drying cycle optimization by interactive selection of the operating condition profiles and simultaneous monitoring of the critical product quality parameters. In order to

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simplify comparison between the S and PS product formulations, most operating conditions were fixed to their usual values (condenser temperature to -65°C , chamber temperature to $+25^{\circ}\text{C}$, total chamber pressure to 10 Pa, final average moisture content target to 0.03 kg/kg) and the only optimized operating condition was the shelf temperature profile.

In all cases, increasing freeze-drying productivity requires increasing the shelf temperature as much as possible, while still satisfying the product stability requirement, i.e. product temperature below the glass transition temperature. Operating the process close to the highest temperature limit is considered beneficial not only for process productivity but also for the distributions of temperature and residual moisture at the end of the drying (19). During the interactive optimisation process, product temperatures are monitored at the bottom of the vial, at the sublimation front and at the top of the product. Glass transition temperatures are computed and displayed for the same points. During primary drying, the limiting factor is usually the bottom temperature. The shelf temperature has to be decreased in time, because the self-cooling effect due to ice sublimation decreases when the mass transfer resistance through the dry layer increases with time. A screenshot of the software tool during the interactive optimisation of the shelf temperature profile is shown in Figure 6.

The top temperature may become limiting, however, if the heat transfer from the chamber is high and the heat conductivity of the dry layer is low, despite the fact that the glass transition temperature at the top increases rapidly when the moisture content of the product decreases. During the secondary drying, the moisture content of the whole product decreases, the glass transition temperature increases everywhere and the shelf temperature can be increased significantly up to a limit imposed by the thermal sensitivity of the biological product.

Variations of the total chamber pressure profile were investigated and were found to have little effect on the product temperature and on the cycle duration. The chamber pressure mainly influences the heat transfer between the shelf and the vial, and similar effect can be easily obtained by manipulating the shelf temperature. Thus maintaining a low pressure and controlling the heat flux by the shelf temperature appears as a good policy (20, 21). It should be noted, however, that in the considered pilot-scale freeze-dryer a significant fraction of the transferred heat (up to 50% at low shelf temperatures) comes from the chamber, mainly by radiation. This need not to be the case in an industrial-scale dryer, when the heat transfer between the shelf and the vials through gas conduction may be dominant and the pressure effect would be stronger.

The best achievable control policies in terms of cycle duration for the two considered product formulations, S and PS, are compared in Table 3. As expected, the PS formulation allows higher shelf temperatures because of its higher glass transition temperature, and hence faster heat transfer resulting in a shorter freeze-drying cycle. In

the considered case, the cycle duration is shorter by 12 hours (33%) which is significant for process scheduling and designing the freeze-drying capacity on an industrial scale.

CONCLUSION

A user-friendly software for interactive selection of the operating conditions in a freeze-drying process was written. It was based on a validated dynamic model of the freeze-drying process, allowing very quick and accurate simulations of the primary and secondary drying stages, as well as of the gradual transition between the primary and secondary drying stages. Process productivity and product quality indicators at critical points were monitored simultaneously during the cycle optimisation process. It was shown that the optimal operating policy as well as the achievable cycle duration strongly depend on the physical properties of the product formulation, namely the glass transition temperature, the sorption isotherm and the desorption kinetic.

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APPENDIX 1. TIME CONSTANTS FOR THE HEAT AND MASS TRANSFER

The considered freeze-drying model assumes quasi steady state heat transfer in the frozen and in the dry product regions, as well as quasi steady state mass transfer in the gas phase of the dry region (Assumption A2). In order to justify these assumptions, the order of magnitude of the involved time constants is estimated below, by considering full dynamic heat and mass balance equations. These equations are based on a lumped parameter approximation (as opposed to a distributed parameter approach, involving partial differential equations), but are sufficient to show that the considered dynamics can be safely neglected compared to the typical duration of a freeze-drying process (100000 s).

Heat transfer dynamics in the frozen product layer

The dynamic heat balance considers heat accumulation in the frozen layer (resulting in temperature variation) and heat fluxes from the shelf and towards the sublimation front:

$$\left(M^{Ice} c^{Ice} + M^{Dry} c^{Dry} \right) \frac{d\Theta^{Frozen}}{dt} = H^{Shelf\ Frozen} (\Theta^{Shelf} - \Theta^{Frozen}) - H^{Frozen\ Front} (\Theta^{Frozen} - \Theta^{Front}) \quad (13)$$

The heat capacity of the frozen layer takes into account the presence of ice and dry matter and depends on the sublimation front position. In the lumped parameter approximation, the equivalent heat transfer coefficient between the shelf and the frozen layer includes heat transfer between the shelf and the product bottom and heat conduction through one half of the frozen layer. Heat transfer between the frozen layer and the sublimation front consists in heat conduction through the other half of the frozen layer. The dynamic heat balance can be written in the following form:

$$\frac{d\Theta^{Frozen}}{dt} = \frac{1}{\tau_{\Theta}^{Shelf\ Frozen}} (\Theta^{Shelf} - \Theta^{Frozen}) - \frac{1}{\tau_{\Theta}^{Frozen\ Front}} (\Theta^{Frozen} - \Theta^{Front}) \quad (14)$$

Substituting typical numerical values, e.g. $Z^{Front} = Z^{Tot} / 2$, $P_t^{Chamb} = 20\text{ Pa}$ etc., one obtains the time constants

$\tau_{\Theta}^{Shelf\ Frozen} = 564\text{ s}$ and $\tau_{\Theta}^{Frozen\ Front} = 4.37\text{ s}$, indicating that the thermal equilibrium between the shelf and the frozen layer is almost achieved in a several minutes, and between the frozen layer and the sublimation front in a few seconds.

Heat transfer dynamics in the dry product layer

A dynamic heat balance for the dry product layer was constructed in a similar way. The heat capacity of the dry layer includes that of the dry product and of the residual moisture and is much smaller than that of the frozen layer. In the lumped parameter formulation, the equivalent heat transfer coefficient between the sublimation front and the dry layer is based on heat conduction through one half of the dry layer. The heat transfer between the dry layer and the chamber includes conduction through the other half of the dry layer and transfer between the product top and the chamber. With typical numerical values, the time constants are $\tau_{\Theta}^{Front\ Dry} = 1.05\text{ s}$ and $\tau_{\Theta}^{Dry\ Chamb} = 8.35\text{ s}$. Thermal equilibrium of the dry product layer is thus achieved in less than one minute.

Mass transfer dynamics in the gaseous phase of the dry product layer

A dynamic mass balance was established for the gaseous phase of the dry product layer. Vapour accumulation in the dry volume was expressed using the perfect gas law. With the lumped parameter formulation, the equivalent mass transfer coefficient between the dry layer and the sublimation front takes into account the transfer resistance of one half of the layer. The mass transfer coefficient between the dry layer and the chamber includes transfer resistance through the other half of the layer and between the product top and the chamber. In order to establish the time constants, the dynamic balance was rewritten in the following form:

$$\frac{dP_w^{Dry}}{dt} = \frac{1}{\tau_p^{Front\ Dry}} (P_w^{Front} - P_w^{Dry}) - \frac{1}{\tau_p^{Dry\ Chamb}} (P_w^{Dry} - P_w^{Chamb}) \quad (15)$$

With typical numerical values, one gets $\tau_p^{Front\ Dry} = 3.9\text{ ms}$ and $\tau_p^{Dry\ Chamb} = 7.2\text{ ms}$. The vapour pressure in the dry product layer reaches quasi steady state quite quickly (milliseconds), as physically expected.

APPENDIX 2. HEAT AND MASS TRANSFER MODEL EQUATIONS

	Primary drying (I)	Secondary drying (II)
Control variables	$\Theta^{Shelf}, \Theta^{Chamb}, \Theta^{Cond}, P_t^{Chamb}$	
State variables	$Z^{Front}, \Theta^{Front}, C^{Bottom}, C^{Front}, C^{Top}$	
Mass transfer		
Mass transfer coefficients	$K^{FrontTop} = k^{FrontTop} \frac{A}{Z^{Tot} - Z^{Front}}$ $K^{TopChamb} = k^{TopChamb} A$ $\frac{1}{K^{FrontCond}} = \frac{1}{K^{FrontTop}} + \frac{1}{K^{TopChamb}} + \frac{1}{K^{ChambCond}}$	
Fixed pressures	$P_w^{Front} = P_w^{Sat}(\Theta^{Front})$	$P_w^{Cond} = P_w^{Sat}(\Theta^{Cond})$ $P_w^{Front} = P_w^{Cond}$
Mass flux	$F^{FrontCond} = K^{FrontCond} (P_w^{Cond} - P_w^{Front})$	
Pressures	$P_w^{Top} = P_w^{Front} + \frac{1}{K^{FrontTop}} F^{FrontCond}$ $P_w^{Chamb} = P_w^{Cond} - \frac{1}{K^{ChambCond}} F^{FrontCond}$	$P_w^{Top} = P_w^{Cond}$ $P_w^{Chamb} = P_w^{Cond}$
Heat transfer		
Heat transfer coefficients	$H^{ShelfBottom} = h^{ShelfBottom} (P_t^{Chamb}, H_2O) A$ $H^{BottomFront} = h^{BottomFront} \frac{A}{Z^{Front}}$ $H^{FrontTop} = h^{FrontTop} \left(\frac{P_w^{Front} + P_w^{Top}}{2}, H_2O \right) \frac{A}{Z^{Tot} - Z^{Front}}$ $H^{TopChamb} = h^{TopChamb} A$ $\frac{1}{H^{ShelfFront}} = \frac{1}{H^{ShelfBottom}} + \frac{1}{H^{BottomFront}}$ $\frac{1}{H^{FrontChamb}} = \frac{1}{H^{FrontTop}} + \frac{1}{H^{TopChamb}}$	$H^{ShelfBottom} = h^{ShelfBottom} (P_t^{Chamb}, N_2) A$ $H^{FrontTop} = h^{FrontTop} \left(\frac{P_w^{Front} + P_w^{Top}}{2}, N_2 \right) \frac{A}{Z^{Tot}}$ $\frac{1}{H^{ShelfChamb}} = \frac{1}{H^{ShelfBottom}} + \frac{1}{H^{FrontTop}} + \frac{1}{H^{TopChamb}}$
Heat fluxes	$Q^{ShelfFront} = H^{ShelfFront} (\Theta^{Front} - \Theta^{Shelf})$ $Q^{FrontChamb} = H^{FrontChamb} (\Theta^{Chamb} - \Theta^{Front})$	$Q^{ShelfChamb} = H^{ShelfChamb} (\Theta^{Chamb} - \Theta^{Shelf})$
Temperatures	$\Theta^{Bottom} = \Theta^{Shelf} + \frac{1}{H^{ShelfBottom}} Q^{ShelfFront}$ $\Theta^{Top} = \Theta^{Front} + \frac{1}{H^{FrontTop}} Q^{FrontChamb}$	$\Theta^{Bottom} = \Theta^{Shelf} + \frac{1}{H^{ShelfBottom}} Q^{ShelfChamb}$ $\Theta^{Front} = \Theta^{Bottom}$
Product quality		
Glass transition	$\Theta_{Glass}^{Bottom} = \Theta_{Glass} (C^{Bottom})$ $\Theta_{Glass}^{Front} = \Theta_{Glass} (C^{Front})$ $\Theta_{Glass}^{Top} = \Theta_{Glass} (C^{Top})$	
Sorption isotherms	$C_{Equ}^{Bottom} = C_{Equ} (1)$ $C_{Equ}^{Front} = C_{Equ} (1)$	$C_{Equ}^{Bottom} = C_{Equ} \left(\frac{P_w^{Bottom}}{P_w^{Sat}(\Theta^{Bottom})} \right)$ $C_{Equ}^{Front} = C_{Equ} \left(\frac{P_w^{Front}}{P_w^{Sat}(\Theta^{Front})} \right)$

	$C_{Equ}^{Top} = C_{Equ} \left(\frac{P_w^{Top}}{P_w^{Sat}(\Theta^{Top})} \right)$	
State evolution equations		
Front position	$\frac{dZ^{Front}}{dt} = \frac{F^{Front\ Cond}}{D_{Ice} A}$	$\frac{dZ^{Front}}{dt} = 0$
Front temperature	$0 = (Q^{Shelf\ Front} - Q^{Front\ Chamb}) - L_{Subl} F^{Front\ Cond}$	$0 = \Theta^{Bottom} - \Theta^{Front}$
Residual water content	$\frac{dC^{Bottom}}{dt} = \frac{1}{\tau_{Des}} (C_{Equ}^{Bottom} - C^{Bottom})$ $\frac{dC^{Front}}{dt} = \frac{1}{\tau_{Des}} (C_{Equ}^{Front} - C^{Front})$ $\frac{dC^{Top}}{dt} = \frac{1}{\tau_{Des}} (C_{Equ}^{Top} - C^{Top})$	
Additional outputs		
Neutral gas press	$P_n^{Chamb} = P_t^{Chamb} - P_w^{Chamb}$	
Average water content	$C^{Ave} = \frac{Z^{Front} (C^{Bottom} + C^{Front}) / 2 + (Z^{Tot} - Z^{Front}) (C^{Front} + C^{Top}) / 2}{Z^{Tot}}$	$C^{Ave} = (C^{Bottom} + C^{Top}) / 2$
Product mass in a vial	$M^{Vial} = M^{Dry} + \frac{C^{Ave}}{1 - C^{Ave}} M^{Dry} + \frac{Z^{Front}}{Z^{Tot}} M^{Ice\ Ini}$	$M^{Vial} = M^{Dry} + \frac{C^{Ave}}{1 - C^{Ave}} M^{Dry}$
Functions		
Sorption isotherm	$C_{Equ}(a_w) = \min \{ C_{Max}, \max \{ C_{Min}, q_{Sorp1}(a_w - q_{Sorp2}) \} \}$	
Transition from primary to secondary drying	$f_{(I)}(Z) = \begin{cases} \sin\left(\frac{\pi Z}{2 Z_{Trans}}\right) & \text{if } Z < Z_{Trans} \\ 1 & \text{if } Z \geq Z_{Trans} \end{cases}$	
Glass transition	$\Theta_{Glass}(C) = \max \left\{ \Theta_{Glass}^{Frozen}, \frac{(1-C)\Theta_{Glass}^{Dry} + q_{Gordon} C \Theta_{Glass}^{Ice}}{(1-C) + q_{Gordon} C} \right\}$	
Heat transfer between shelf and product bottom	$h^{Shelf\ Bottom}(P, gas) = h_{Contact+Rad}^{Shelf\ Bottom} + q_{gas} h_{Cond}^{Shelf\ Bottom} \frac{P}{1 + P / P_{Trans}^{Shelf\ Bottom}}$	
Heat transfer between sublimation front and product top	$h^{Front\ Top}(P, gas) = h_{Walls+Rad}^{Front\ Top} + q_{gas} h_{Cond}^{Front\ Top} \frac{P}{1 + P / P_{Trans}^{Front\ Top}}$	
Saturated vapour pressure	$P_w^{Sat}(\Theta) = \exp\left(-\frac{6139.6}{\Theta + 273.15} + 28.8912\right)$	

REFERENCES

1. Leuenberger, H., M. Plitzko and M. Puchkov. 2006. Spray freeze drying in a fluidized bed at normal and low pressure. *Drying Technology* 24: 711-719.
2. Pikal, M. J. 1985. Use of a laboratory data in freeze drying process design : heat and mass transfer coefficients and the computer simulation of freeze drying. *Journal of Parenter Science and Technology* 39: 115-138.

3. Chang, B. S. and N. L. Fisher. 1995. Development of an efficient single-step freeze-drying cycle for protein formulations. *Pharmaceutical Research* 12: 831-837.
4. Tang, X. L. and M. J. Pikal. 2004. Design of freeze-drying processes for pharmaceuticals: practical advice. *Pharmaceutical Research* 21: 191-200.
5. Pikal, M. J. and S. Shah. 1990. The collapse temperature in freeze-drying: dependence on measurement methodology and rate of water removal from the glassy state. *International Journal of Pharmacy* 62: 165-186.
6. Alves, O. and Y. H. Roos. 2006. Advances in multi-purpose drying operations with phase and state transitions. *Drying Technology* 24: 383-396.
7. Chouvinc, P., S. Vessot, J. Andrieu and P. Vacus. 2004. Optimization of the freeze-drying cycle: A new model for pressure rise analysis. *Drying Technology* 22: 1577-1601.
8. Sadikoglu, H. 2005. Optimal control of the secondary drying stage of freeze drying of solutions in vials using variational calculus. *Drying Technology* 23: 33-57.
9. Sadikoglu, H., M. Ozdemir and M. Seker. 2003. Optimal control of the primary drying stage of freeze drying of solutions in vials using variational calculus. *Drying Technology* 21: 1307-1331.
10. Nam, J. H. and C. S. Song. 2005. An efficient calculation of multidimensional freeze-drying problems using fixed grid method. *Drying Technology* 23: 2491-2511.
11. Hottot, A., R. Peczalski, S. Vessot and J. Andrieu. 2006. Freeze-drying of pharmaceutical proteins in vials: Modeling of freezing and sublimation steps. *Drying Technology* 24: 561-570.
12. Song, C. S., J. H. Nam, C. J. Kim and S. T. Ro. 2002. A finite volume analysis of vacuum freeze drying process of skim milk solution in trays and vials. *Drying Technology* 20: 283-305.
13. Mascarenhas, W. J., H. U. Akay and M. J. Pikal. 1997. A computational model for finite element analysis of the freeze-drying process. *Computer Methods in Applied Mechanics and Engineering* 148: 105-124.
14. Passot, S., F. Fonseca, M. Alarcon-Lorca, D. Rolland and M. Marin. 2005. Physical characterisation of formulations for the development of two stable freeze-dried proteins during both dried and liquid storage. *European Journal of Pharmaceutics and Biopharmaceutics* 60: 335-348.

15. Sadikoglu, H. and A. I. Liapis. 1997. Mathematical modelling of the primary and secondary drying stages of bulk solution freeze-drying in trays: parameter estimation and model discrimination by comparison of theoretical results with experimental data. *Drying Technology* 15: 791-810.
16. Nastaj, J. F. and B. Ambrozek. 2005. Modeling of vacuum desorption in freeze-drying process. *Drying Technology* 23: 1693-1709.
17. Gan, K. H., O. K. Crosser and A. I. Liapis. 2005. Lyophilization in vials on trays: Effects of tray side. *Drying Technology* 23: 341-363.
18. Hua, Z. Z., B. G. Li, Z. J. Liu and D. W. Sun. 2003. Freeze-drying of liposomes with cryoprotectants and its effect on retention rate of encapsulated ftorafur and vitamin A. *Drying Technology* 21: 1491-1505.
19. Gan, K. H., R. Bruttini, O. K. Crosser and A. I. Liapis. 2004. Heating policies during the primary and secondary drying stages of the lyophilization process in vials: effects of the arrangement of vials in clusters of square and hexagonal arrays on trays. *Drying Technology* 22: 1539-1575.
20. Adams, G. D. J. and J. R. Ramsay. 1996. Optimizing the lyophilization cycle and the consequences of collapse on the pharmaceutical acceptability of Erwinia L-Asparaginase. *Journal of Pharmaceutical Sciences* 85: 1301-1305.
21. Ronzi, E., A. Capalongo, G. Rovero, E. Bucci, S. Mondini and A. Falbo. 2003. Optimisation of a freeze-drying process of high purity Factor VIII and Factor IX concentrates. *Chemical Engineering and Processing* 42: 751-757.

Nomenclature

Symbol	Units	Significance
A	m ²	Product cross area
a _w	Pa / Pa	Water activity
C	kg / kg	Moisture content, wet basis
D	kg / m ³	Density
F	kg / s	Mass flux
H	W / K	Heat transfer coefficient
h	W / (K m)	Heat conductivity
h	W / (K m ²)	Unitary heat transfer coefficient
K	kg / (s Pa)	Mass transfer coefficient
k	kg / (s Pa m)	Mass conductivity
k	kg / (s Pa m ²)	Unitary mass transfer coefficient
L _{Subl}	J / kg	Specific sublimation heat
M	kg	Mass for one vial
N		Number
P	Pa	Pressure
Q	W	Heat flux
q		Empirical model coefficient
R	m	Radius
Z	m	Position (height) in the product layer (origin = bottom)
Θ	°C	Temperature
τ	s	Time constant

Upper indices

Ave	Average
Bottom	Product bottom
Chamb	Chamber
Cond	Condenser
Dry	Dry product layer
Front	Sublimation front
Frozen	Frozen product layer
Ice	Ice
Sat	Saturation
Shelf	Temperature controlled shelf
Top	Product top
Tot	Total
Vial	Product vial

Lower indices

(I)	Primary drying stage
(II)	Secondary drying stage
Coll	Collapse
Cond	Conduction
Contact	By contact
Des	Desorption
Equ	Equilibrium
gas	Dominant gas composition: either water vapour (H ₂ O) or nitrogen (N ₂)
Glass	Glass transition
Gordon	Gordon-Taylor formula for glass transition temperature
Ini	Initial
Max	Maximum
Min	Minimum
n	Neutral gas (nitrogen)
Rad	By radiation
Sorp	Sorption isotherm
t	Total
Trans	Transition; either from primary to secondary drying or from low pressure (molecular) to high pressure regime
w	Water vapour
Walls	Conduction through vial walls

Table 1. Formulation-specific model parameters

	Glass transition			Sorption and desorption					Mass transfer
	Θ_{Glass}^{Frozen}	Θ_{Glass}^{Dry}	q_{Gordon}	C_{Min}	C_{Max}	q_{Sorp1}	q_{Sorp2}	τ_{Des}	$k^{FrontTop}$
	°C	°C		kg/kg	kg/kg			s	kg/(s m Pa)
S	-34	65	5.4	0	0.153	0.177	-0.026	12600	$12 \cdot 10^{-9}$
PS	-28	124	8.5	0.0167	0.144	0.459	0.188	10600	$12 \cdot 10^{-9}$

Table 2. Formulation-independent model parameters

Vials and filling						Ice properties			
N^{Vial}	R^{Vial}	M^{Dry}	M_{Ini}^{Ice}	Z^{Tot}	Z^{Trans}	Θ_{Glass}^{Ice}	D^{Ice}	L_{Subl}	$h^{BottomFront}$
	m	kg	kg	m	m	°C	kg/m ³	J/kg	W/(m K)
213	$7.12 \cdot 10^{-3}$	$0.05 \cdot 10^{-3}$	$0.95 \cdot 10^{-3}$	$6.82 \cdot 10^{-3}$	$1.70 \cdot 10^{-3}$	-135	920	$2.83 \cdot 10^6$	2.4

Mass transfer			Heat transfer between shelf and product bottom				
$k^{FrontTop}$	$k^{TopChamb}$	$K^{ChambConc}$	$h_{Contact+Rad}^{ShelfBottom}$	$q_{H_2O}^{ShelfBottom}$	$q_{N_2}^{ShelfBottom}$	$h_{Cond}^{ShelfBottom}$	$P_{Trans}^{ShelfBottom}$
kg/(s m Pa)	kg/(s m ² Pa)	kg/(s Pa)	W/(m ² K)			W/(m ² K Pa)	Pa
$1.2 \cdot 10^{-8}$	$8 \cdot 10^{-6}$	$5 \cdot 10^{-7}$	1	1	0.625	0.6	100

Heat transfer between sublimation front and product top					Heat transfer between product top and chamber
$h_{Walls+Rad}^{FrontTop}$	$q_{H_2O}^{FrontTop}$	$q_{N_2}^{FrontTop}$	$h_{Cond}^{FrontTop}$	$P_{Trans}^{FrontTop}$	$h^{TopChamb}$
W/(m K)			W/(m K Pa)	Pa	W/(m ² K)
$50 \cdot 10^{-3}$	1	0.813	$327 \cdot 10^{-6}$	100	5

Interactive freeze-drying optimisation

Table 3. Optimisation results for the shelf temperature profile

Formulation	Initial (maximum) shelf temperature in primary drying °C	Final (minimum) shelf temperature in primary drying °C	Total cycle duration h
S	-29	-48	36
PS	+12	-30	24

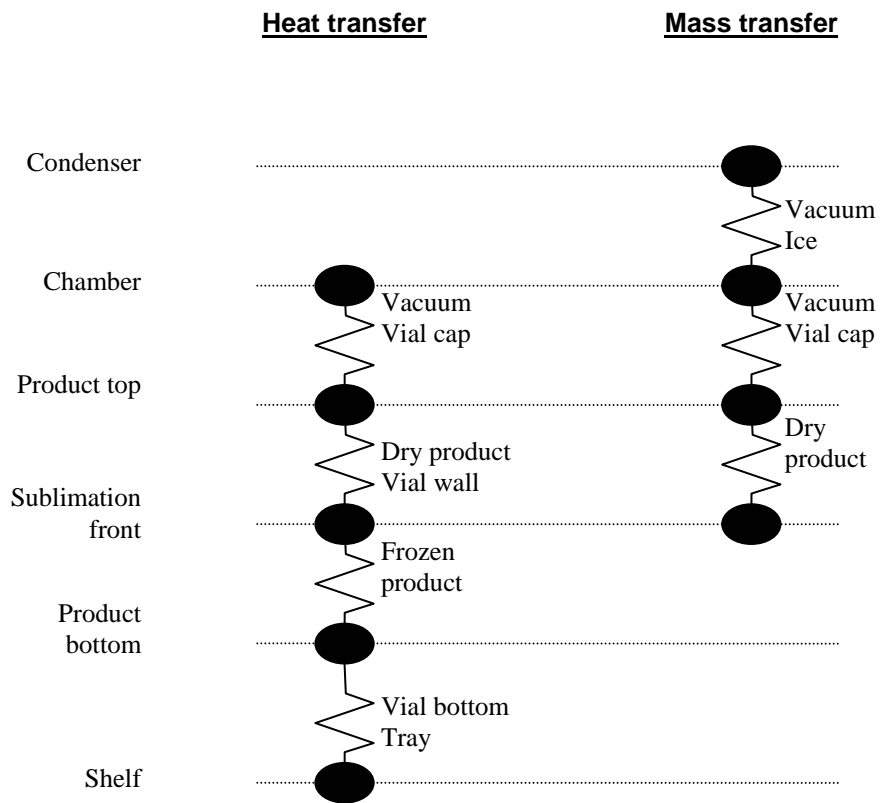


Figure 1. Schematic representation of heat and mass transfers between the characteristic points in the product and in the freeze-dryer.

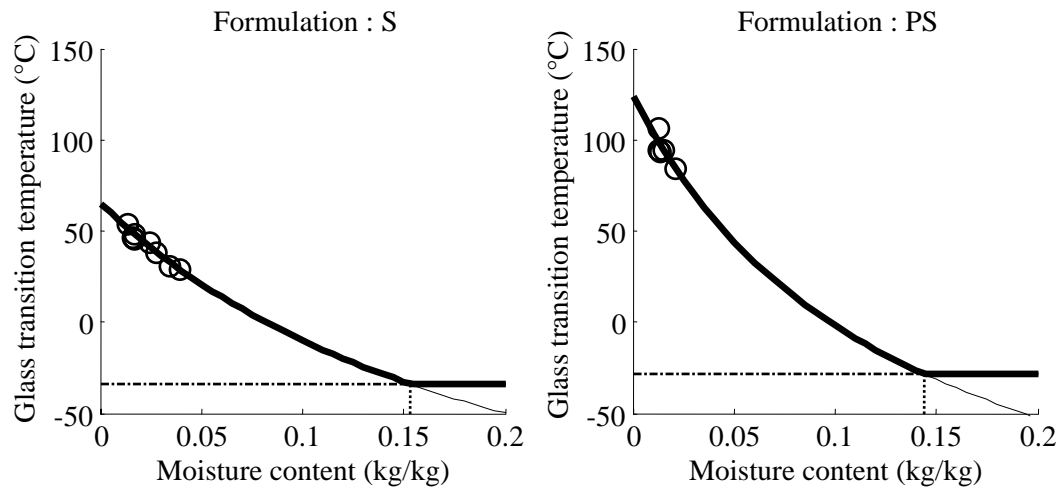


Figure 2. Glass transition temperature models. Measured values in dry product (o), measured value in frozen product (---), Gordon-Taylor model (—), maximum moisture content of the freeze-concentrated product (.....).

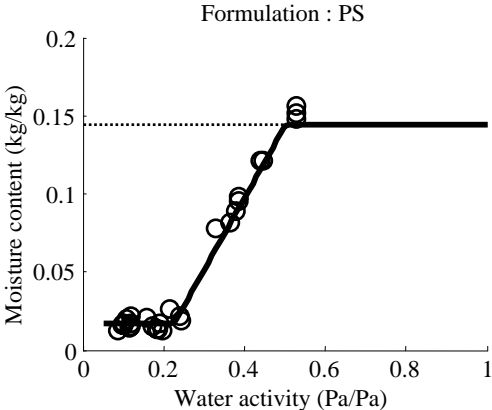


Figure 3. Sorption isotherm models. Measured values (o), piecewise linear model (—), maximum moisture content of the freeze-concentrated product (.....).

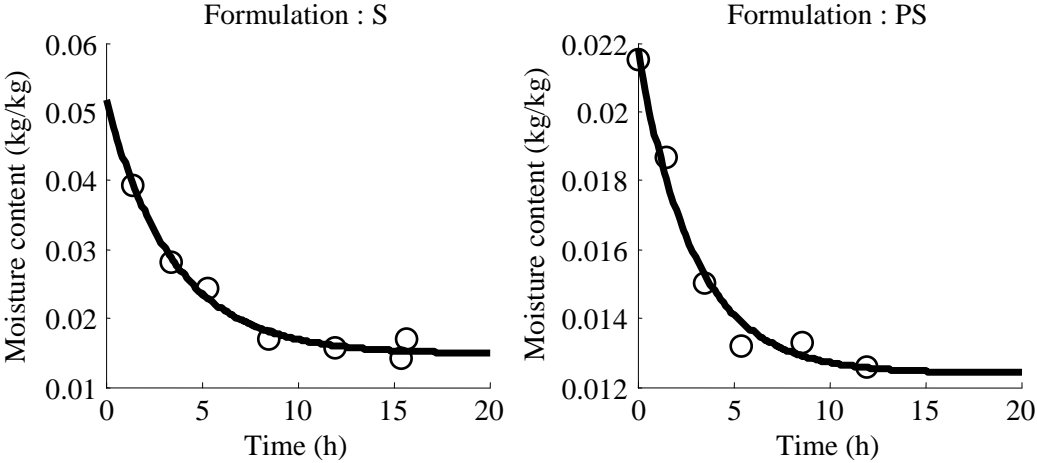


Figure 4. Desorption kinetics. Measured values (o), exponential first order rate model (-).

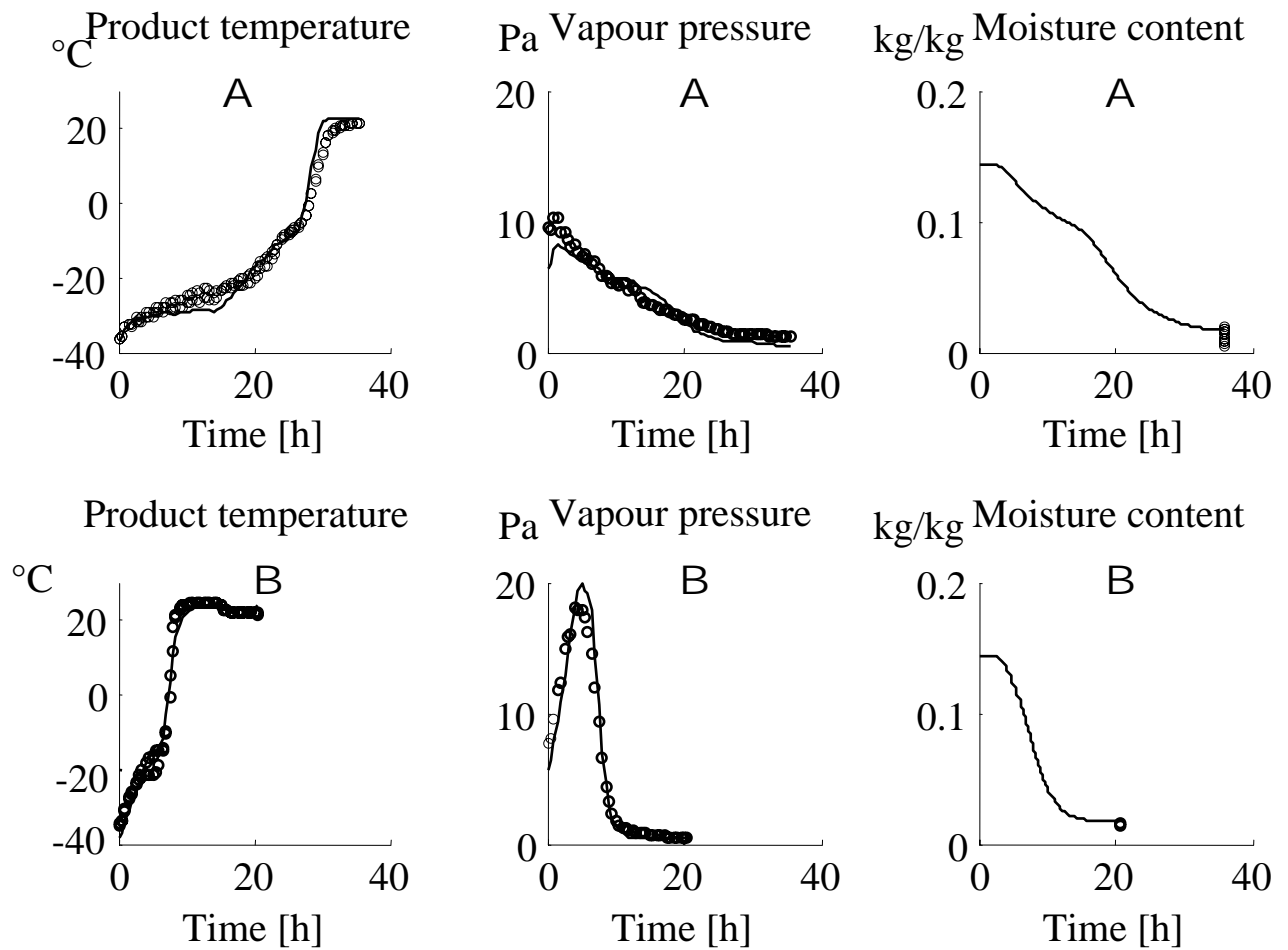


Figure 5. Dynamic model validation. Formulation: PS. **A:** Shelf temperature at -25°C and total chamber pressure at 10 Pa during primary drying. **B:** Shelf temperature at +25°C and total chamber pressure at 34 Pa during primary drying. Measured values (o) and model predictions (-).

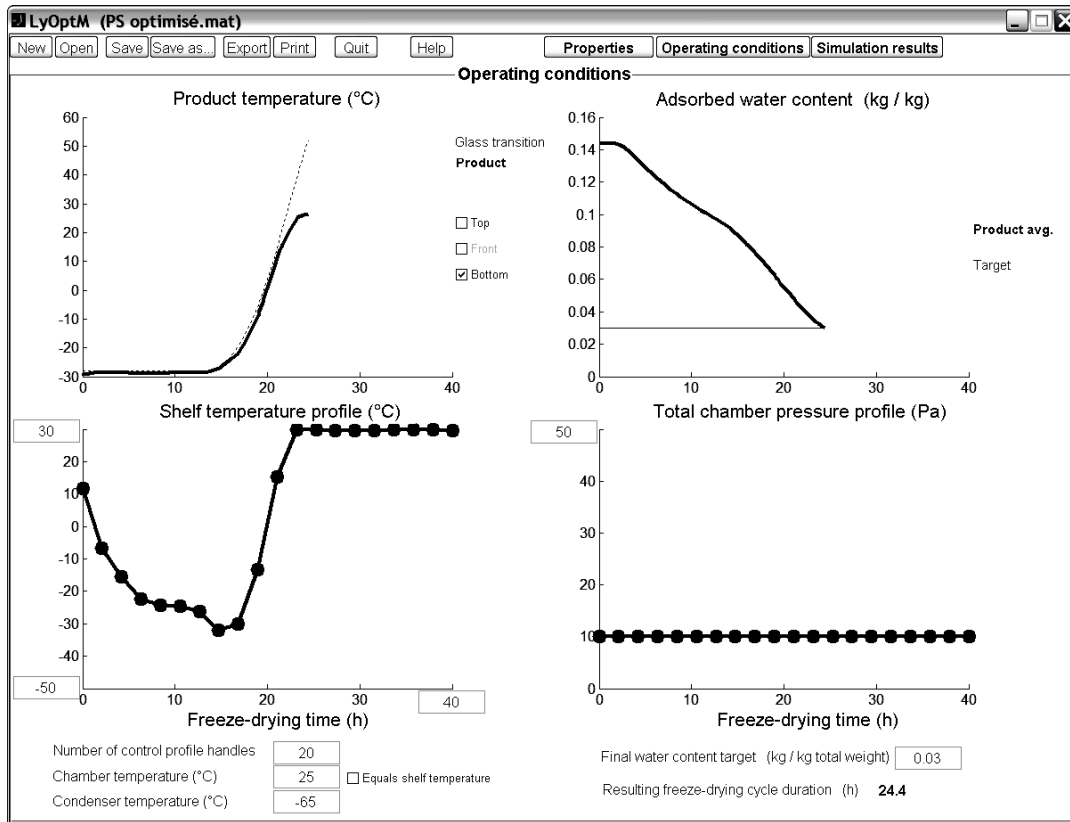


Figure 6. Screenshot of the freeze-drying cycle optimisation tool. Interactive selection of the operating conditions for the PS formulation.