Use of Physiologically-Based Kinetics Modelling to Reliably Predict Internal Concentrations of the UV Filter, Homosalate, After Repeated Oral and Topical Application

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Ethical and legal considerations have led to increased use of non-animal methods to evaluate the safety of chemicals for human use. We describe the development and qualification of a physiologically-based kinetics (PBK) model for the cosmetic UV filter ingredient, homosalate, to support its safety without the need of generating further animal data. The intravenous (IV) rat PBK model, using PK-Sim®, was developed and validated using legacy in vivo data generated prior to the 2013 EU animal-testing ban. Input data included literature or predicted physicochemical and pharmacokinetic properties. The refined IV rat PBK model was subject to sensitivity analysis to identify homosalate-specific sensitive parameters impacting the prediction of Cmax (more sensitive than AUC(0–∞)). These were then considered, together with population modeling, to calculate the confidence interval (CI) 95% Cmax and AUC(0–∞). Final model parameters were established by visual inspection of the simulations and biological plausibility. The IV rat model was extrapolated to oral administration, and used to estimate internal exposures to doses tested in an oral repeated dose toxicity study. Next, a human PBK dermal model was developed using measured human in vitro ADME data and a module to represent the dermal route. Model performance was confirmed by comparing predicted and measured values from a US-FDA clinical trial (Identifier: NCT03582215, https://clinicaltrials.gov/). Final exposure estimations were obtained in a virtual population and considering the in vitro and input parameter uncertainty. This model was then used to estimate the Cmax and AUC(0–24 h) of homosalate according to consumer use in a sunscreen. The developed rat and human PBK models had a good biological basis and reproduced in vivo legacy rat and human clinical kinetics data. They also complied with the most recent WHO and OECD

Abbreviations: ADME, absorption, distribution, metabolism and excretion; BSA, body surface area; Css, concentration at steady state; CI, confidence interval; CLs, total plasma clearance; ECCS, Extended Clearance Classification System; Fa, oral absorption; Fb, oral bioavailability; GFR, glomerular filtration rate; IV, intravenous; MoIE, Margin of Internal Exposure; NOAEL, no observed adverse effect level; OECD, Organization for Economic Co-operation and Development; PBK, physiologically-based kinetics; PoD, point of departure; SCCS, Scientific Committee on Consumer Safety; SVM, Support Vector Machine; TK/PK, toxico/pharmacokinetics; WHO, World Health Organization.
recommendations for assessing the confidence level. In conclusion, we have developed a PBK model which predicted reasonably well the internal exposure of homosalate according to different exposure scenarios with a medium to high level of confidence. In the absence of in vivo data, such human PBK models will be the heart of future completely non-animal risk assessments; therefore, valid approaches will be key in gaining their regulatory acceptance.

Clinical Trial Registration: https://clinicaltrials.gov/, identifier, NCT03582215

Keywords: homosalate, physiologically-based kinetics models, UV filter, plasma concentration, dermal application

INTRODUCTION

All chemicals should be assessed for potential toxicity to humans before they are used in products. Traditionally, this has been achieved using standardized animal studies for evaluation of local and systemic effects. These are used to identify adverse effects, target organs, as well as the dose below which no adverse effects are observed (the no observed adverse effect level [NOAEL]). The latter is used as a point of departure (PoD) which can be compared with dose levels experienced by humans through use of the product. However, ethical and legal considerations have led to a change in paradigm towards non-animal methods to evaluate the safety of chemicals for human use. This change is particularly important for cosmetics ingredients, for which animal testing has been banned since March 2013 (Eu, 2009), although non-animal approaches are increasingly of interest for other sectors, such as industrial chemicals and environmental contaminants (Thomas et al., 2019).

The availability of toxico/pharmacokinetics (TK/PK) data greatly increases confidence in risk assessment decisions and can also be valuable in supporting the replacement, reduction and refinement of animal use (3Rs) (Creton et al., 2009). In the absence of animal or human kinetics data, the extrapolation of external doses (in mg/kg/day) to internal dose metrics ($C_{\text{max}}$, AUC or concentration at steady state ($C_{\text{ss}}$)) must be predicted. Therefore, physiologically-based kinetics (PBK) models are important tools in safety assessment. These are mathematical models used to quantify and predict the absorption, distribution, metabolism and excretion of a chemical following exposure. They are composed of interconnected compartments representing various tissues/organs described by mass balance differential equations that are solved to predict the amount of chemical in each compartment over time (Gerlowski and Jain, 1983). PBK models typically rely on three types of parameters; physiological (e.g., tissue volumes, blood flows), physicochemical (e.g., octanol:water partitioning, vapor pressure, water solubility), and biochemical (e.g., absorption rates, metabolism, clearances). The necessary physiological parameters for several species (mouse, rat, dog and human) are available in the literature (Brown et al., 1997). Tissue:blood partition coefficients for a chemical can be estimated using quantitative structure-property relationships (Peyret et al., 2010), while the clearance of the chemical in different species can be determined using in vitro studies with hepatocytes or cellular fractions and incorporated into the PBK model using in vitro to in vivo extrapolation (Yoon et al., 2012). This modelling approach allows internal concentrations resulting from external exposures to be predicted, allowing comparisons including across species and exposure routes (Clewell and Andersen, 1985).

Guidelines for the application, use, and reporting of PBK models for drugs and chemicals have been published by the FDA and EMA (Ema, 2016; Us fda, 2018). Guidance documents are also available from the Organization for Economic Cooperation and Development (OECD) and World Health Organization (WHO) for the characterization and application of PBK models in the risk assessment of chemicals (WHO, 2010; OECD, 2021), with the aim to harmonize and facilitate their use by chemical developers and regulators. This is particularly important for the cosmetics industry, which fully supports the animal testing ban under the European Union Cosmetics Products Regulation and has decades of commitment in promoting the use of alternatives to animal testing for safety assessment of cosmetics. Indeed, the ambition of Cosmetics Europe (the European cosmetics industry trade association, https://cosmetics-europe.eu/) is to develop non-animal tools and approaches (Desprez et al., 2018). However, historical pre-ban animal toxicology data exist for many cosmetic ingredients, and an important principle in risk assessment is that all available relevant data need to be taken into account when arriving at a safety decision. Such historical data exist for the UV filter homosalate, an ester of salicylic acid and a key cosmetic ingredient in sunscreens. This ingredient provides consumers with protection from the Sun’s [UVA/B] exposure and thus plays a key role in the protection against skin cancer caused by UV exposure. Here, we describe the development and qualification of PBK models for homosalate, which were built to put exposures in an historical animal study into context and substantiate the human safety evaluation without generating further animal data. The aim of this work was to use the PBK model of homosalate to perform route-to-route and interspecies extrapolations to translate both the oral exposures from a historical rat study, and dermal exposures in consumers using sunscreen to internal dose metrics. This could help replace default assessment factors with more specific, substance-derived factors and exclude or refine a toxicokinetic factor in the calculation of margins of safety (SCCS, 2021).
TABLE 1 | Physicochemical properties of homosalate [taken from the REACH dossier (ECHA 2020)].

| Property                          | Value                                      |
|-----------------------------------|--------------------------------------------|
| Log P<sub>OW</sub>                | 6.34 at 40°C                               |
| Boiling point                     | 295.1°C at 101.3 kPa                       |
| Melting point                     | < -20°C at 101.3 kPa                       |
| Vapor pressure                    | 0.015 Pa at 25°C                           |
| Molecular weight                  | 262.344                                    |
| Water solubility                  | 0.4 mg/L at 25°C                           |
| pKa                               | 8.1 ± 0.3 at 20°C                          |
| Relative density at 20°C relative to water at 4°C | 1.0512 (1.050–1.053) |

METHODS

Step 1: Development of the Rat PBK Model

Physiological and Physicochemical Properties

Table 1 displays the information on the physicochemical properties of homosalate. The rat physiological parameters were scaled from the mean body weight of 0.23 kg, age 40 weeks, and a glomerular filtration rate (GFR) of 57 ml/min/100 g organ (Davies and Morris, 1993).

Parameter Identification

The base mean model in rats was built using animal data to identify an appropriate structure to describe the plasma kinetics of a single intravenous (IV) bolus dose of 0.5 mg/kg homosalate reported by Kim et al. (Kim et al., 2014). WebPlotDigitizer-4.3 software (https://automeris.io/WebPlotDigitizer/) was used to extract data from plots presented in the paper. Unknown parameters were identified using the Parameter Identification Module in PK-Sim® and using the Monte-Carlo algorithm. The parameter identification function in PK-Sim was used to optimize the model input parameters, using available kinetics data. The final model parameters were established by visual inspection of the resulting description of data and biological plausibility.

The parameters used for rat model building are summarized in Table 2. The distribution process of homosalate was according to the partition coefficient calculation by Rodgers and Rowland (Rodgers and Rowland, 2006) and the cellular permeability using a charge-dependent model (Schmitt, 2008). Rodgers and Rowland developed two models: one for moderate to strong bases (pKa > 7) and another model for weak acids and neutral drugs (pKa < 7). The predictions of the steady-state unbound tissue-plasma water partition coefficient, which considers the partitioning of the drug into neutral lipids and phospholipids, the dissolution into tissue water, and the electrostatic interactions with tissue phospholipids. The permeability model developed by Schmitt et al. is calculated from the physicochemical properties and considered the degree of dissociation of acids and bases assuming that the permeabilities for charged species are significantly smaller than for neutral species.

These were selected after testing the available organ-plasma partition coefficient and cell permeability calculation methods built in PK-Sim 9.1. Homosalate fraction unbound (Fu) was initially set at 2.0 ± 0.2%, which was determined from in silico and in vitro measurements. The contribution of transporters to the distribution of homosalate was evaluated using two in silico tools: a substrate classification model provided by SimulationPlus and SwissADME (Daina et al., 2017). These indicated (with a high probability) that homosalate was not a transporter substrate (see Supplementary Materials); therefore, the

TABLE 2 | Homosalate parameters used for rat and human model building. Values were obtained from human based in vitro assays from the literature, collected from the SCCS report on homosalate 2007 (SCCP, 2007) or newly generated data (unpublished). NA = not applicable.

| Property                              | Rat                                         | Human                                      |
|---------------------------------------|---------------------------------------------|--------------------------------------------|
| Dermal bioavailability                |                                             |                                             |
| SCOS report                           | NA                                          | 2% (SCCP, 2007)                           |
| CRL Dermal penetration                | NA                                          | 3.86% (Finlayson, 2021)                    |
| CRL Dermal penetration plus 1 SD     | NA                                          | 5.3% (Finlayson, 2021)                     |
| Oral bioavailability                  |                                             |                                             |
| A Conservative approach              | 50% (assumption, SCCS default) (adjusted Fa = 81%) |                                             |
| Model-based estimation                | Fa = 100%, Fb = 83%                         | NA                                         |
| Distribution                          |                                             |                                             |
| Tissue:plasma partition coefficient   |                                             |                                             |
| Cellular permeabilities               | Rodgers and Rowland (Rodgers and Rowland, 2006) | Rodgers and Rowland (Rodgers and Rowland, 2006) |
| Fraction unbound (Fu)                 | Charge dependent (Schmitt, 2008)            | Charge dependent (Schmitt, 2008)           |
| Elimination                           |                                             |                                             |
| Total plasma clearance (Cl<sub>t</sub>) | 6 L/h/kg (Kim et al., 2014)                 | 59.6 ± 2.7 μL min<sup>-1</sup>.10<sup>6</sup> cells<sup>-1</sup> |
| Cl<sub>t</sub><sub>h,pkw</sub> (primary human hepatocytes) | NA                                          | 11.64 ± 0.53 min                          |
| In vitro half-life (primary human hepatocytes) | NA                                          | No (in silico)                            |
| Transporters Substrate                | No (in silico)                             | No (in silico)                            |
contribution of these was excluded from the model. Since the focus of this work was on the parent compound, none of the homosalate metabolites were modeled. The elimination rate, represented as total plasma clearance (CLs) of 6 L/h/kg, was taken from the in vivo rat study (Kim et al., 2014).

Sensitivity Analysis, and Uncertainty Calculations
The refined IV rat PBK model was subject to sensitivity analysis relating to organism- and homosalate-specific input parameters using PK-Sim, which identifies a set of variables that impact the estimated kinetics. The sensitivity analysis was conducted, according to PK-Sim (PK-Sim Sensitivity Analysis, 2021) in five steps, with a variation range of 10%/step. The sensitivity for the Pharmacokinetics Parameter = PKj to an input parameter = [pi] was then calculated as the ratio of the relative change of that Pharmacokinetics Parameter [ = (ΔPKj)/PKj ] and the relative variation of the input parameter [ = (Δpi)/pi ]:

\[
S_{ij} = \frac{\Delta PK_j}{\Delta Pi} \cdot \frac{Pi}{PK_j}
\]

The sensitivities are dimensionless quantities calculated as the average of several sensitivities based on different variations. Sensitivity analysis results are presented, with sensitivities according to the WHO guidelines (WHO, 2010) as high (absolute value ≥ 0.5), medium (absolute value ≥ 0.2 but less than 0.5) or low (absolute value ≥ 0.1 but < 0.2); parameters with sensitivities < 0.1 are not listed.

Uncertainty expressed by the standard deviation (SD) of the kinetics parameters was considered in the confidence interval CI (5–95)%, such that the influencing parameters were ranked by the SD around the mean of the homosalate-specific parameters. The uncertainty of the in vitro generated inputs was also based on the SD. The variations in the physiology and anatomy were considered through the population modeling, where a virtual population was built to scale those parameters.

Uncertainty of the kinetics parameters was expressed as the confidence interval CI (5–95)%, such that the influencing parameters were ranked by the standard deviation (SD) around the mean of the homosalate-specific parameters. The uncertainty of the in vitro generated inputs was also based on the SD. The variations in the physiology and anatomy were considered through the population modeling.

To evaluate the performance of the refined model, it was also used to estimate the plasma concentration of the additional IV doses reported in the study by Kim et al. (Kim et al., 2014).

Development of the Oral Route Rat PBK Model and Performance
The IV rat model was extrapolated to oral administration in order to convert the doses of 60, 120, 300, and 750 mg/kg/day tested in a repeated dose toxicity study in rats (conducted before March 2013, reported in the Scientific Committee on Consumer Safety (SCCS) dossier (SCCS, 2020)) to an internal plasma concentration. The parameters used for rat model building are summarized in Table 2. A model-based approach was used to calculate the intestinal permeability coefficients based on the molecular weight and membrane affinity (Thelen et al., 2011), implemented in PK-Sim 9.1. The refined oral rat PBK model was subject to sensitivity analysis as described above. The oral bioavailability (Fa) was estimated using in silico modelling and was predicted to be almost complete, while, the oral bioavailability (Fb) was predicted to be 83%, which might be lower than Fa due to the first-pass effect of liver metabolism.

Doses tested in a repeated-dose study according to the OECD 422 Test Guideline were simulated to estimate the Cmax and the area under the curve (AUC0–24) for each dose. AUC0–24 values were estimated on the last dosing day (Day 49). The uncertainty of the influencing parameters, including the calculated intestinal permeability, was evaluated according to the Cmax CI (5–95)%.

Step 2: Development of the Human PBK Dermal Model
An interspecies extrapolation was conducted by developing the rat model to a human PBK dermal model. The oral administration is not relevant for the human PBK model; therefore, only dermal exposure was considered as a route of exposure.

Physiological Properties and Spatial Structure
The spatial structures of the human and rat whole-body models are almost identical, apart from two additional compartments (organs: saliva and gallbladder) in the human model. The anthropometric (height, weight) and physiological parameters in human-adults and rats (e.g., blood flow, organ volumes, binding protein concentrations, hematocrit, cardiac output) were from the literature (Mordenti, 1986; Davies and Morris, 1993; Edginton et al., 2006) or were default values in PK-Sim®. The European human individual was characterized by several parameters representing the mean values of age (30 years), body weight (60 kg), and height (163 cm), BMI (22.58 kg/m²), body surface area (BSA, 1.65 m²) and GFR (107.44 ml/min) (taken from Valentin (2002)). The parameters used for human model building are summarized in Table 2. The same mathematical equations for cellular permeability and Fu established in the rat PBK model were used for the human PBK model. The liver was assumed to be the only site of metabolism and the elimination rate of 59.6 ± 2.7 µL min⁻¹·10⁶ hepatocytes⁻¹ was extrapolated to total hepatic clearance using the IVIVE calculation in PK-Sim 9.1. Based on the Extended Clearance Classification System (ECCS) classification (Varma et al., 2015) and the high plasma binding of homosalate, renal clearance was considered to be negligible and thus was set to zero as a conservative assumption.

Dermal Model
The generic human whole-body model was extended with a dermal model for dermal delivery published by (Dancik et al., 2013). This model assumes the skin is a multilayered slab into which compounds can diffuse. Each slab layer corresponds to a skin layer (stratum corneum, viable epidermis and dermis), each of which exhibit diffusion parameters according to its physical and chemical properties. The skin model was built in MoBi (Bayer Technology Services, Leverkusen, Germany (Hamadeh...
et al., 2019). The skin permeation model has an air compartment, surface pool compartments, vehicle compartment, skin compartment, and in vivo link compartment. The skin compartment consists of three sub-compartments representing the skin layers: stratum corneum sub-compartment, epidermis sub-compartment, dermis sub-compartment. Each of these skin layer sub-compartments comprises four sub-sub-compartments, each of which is composed of ten sub-layers. Dermal clearance in in vivo simulations of homosalate undergoes passive transport from all sub-layers of the skin layers into the bloodstream. The diffusion of homosalate through the skin sub-layers was simulated using the methods of finite differences and Fick’s law (Fick, 1855).

Several in vitro studies reported dermal penetration of homosalate in frozen non-viable human skin ranged from 1.4 to 3.86%, which was assumed to be due to differences in the formulations used. In these studies, dermal penetration was taken from the SCCS opinion (2% (SCCS, 2020)) and from Finlayson (a mean of 3.86% and the mean plus one SD, 5.3%) to estimate the kinetics parameters following the dermal exposure. Several dermal models were built to simulate the different dermal penetration obtained by the in vitro investigations. The main parameters used in the dermal model are listed in the Supplementary Materials. Although in vitro assays indicated first-pass metabolism of homosalate in the skin (data not shown), this was excluded from the PBK modeling of dermal exposure. The model assumed that the total penetrated measured amount was homosalate and no metabolites, as a worst-case scenario for systemic exposure to homosalate.

Model Performance
The PBK model performance was based on a comparison with results from a clinical trial (Matta et al., 2020). In this study, three US commercially available sunscreen formulations containing 10–15% homosalate and other UV filters under maximal use conditions were topically applied to 12 human volunteers. The sunscreens were applied once on Day 1 and then 4 times (with 24-h intervals) on Day 2 through Day 4, and the plasma concentrations were measured at various time points. The non-aerosol spray containing 10% homosalate was selected as a more conservative comparison compared to the pump spray since it resulted in higher plasma concentrations (17.9 ng/ml compared to 13.9 ng/ml, respectively). Homosalate pump spray was applied at 2 mg/cm² to 75% of the body surface area. The actual homosalate dose applied to 19000 cm² was 150 μg/cm².

Step 3: Simulation of Kinetics Following Administration to Rats and Human Virtual Populations
The developed PBK models were used to simulate plasma concentration to homosalate after repeated oral exposure of all the doses administered to rats and after repeated dermal as sunscreen lotion applied on the whole body of humans. The oral dosing to rats was according to an OECD 422 (OECD, 1996) compliant repeated dose toxicity study, in which 20, 60 and 120 mg/kg homosalate was administered daily for 49 days according to the design of the OECD 422 compliant study conducted prior to March 2013 and reported in the REACH dossier (ECHA, 2020; SCCS, 2020). The virtual rat population was for 100 rats, which was generated from the individual mean value by scaling the body weight from 0.185 to 0.275 kg.

In human studies, sunscreen containing 10% homosalate was topically applied to 17500 cm² of skin twice a day (18 g per day, equivalent to 103 μg/cm²/day, according to the typical usage reported by the SCCS (SCCS, 2021)). The maximum plasma concentrations at CI95% were simulated in a virtual population and considering the uncertainty of the in vitro and input parameters. The PBK human model was built for individuals, with the mean values for anatomy and physiology from the European population. Therefore, population PBK modeling was applied to 100 individuals to consider the variations of the individual anatomy and physiology, and the SD of the homosalate-specific sensitive parameters. The age-based population was generated using PK-Sim by scaling the age from 16 to 70, where the body weight ranged from 45 to 100 kg. The corresponding physiology and anatomy parameters were dependently ranged. The normal distribution of input parameters was implemented, along with the Monte-Carlo algorithm for a random-sampling method. The simulation duration was up to 30 days of bi-daily dermal administration. ADME parameters used for human model building are summarized in Table 2. These included intrinsic clearance and half-life values calculated from incubations with cryopreserved primary human hepatocyte (PHH) suspensions over 2-h (according to the method described by Eilstein et al. (2020)). It was not possible to measure plasma protein binding using standard methods such as rapid equilibrium dialysis, ultrafiltration or ultracentrifugation due to the high non-specific binding to plastic and the hydrolysis of homosalate in plasma (data not shown). Therefore, dedicated protocol was used to overcome these issues, whereby plasma protein binding was calculated from a series of incubations of skin S9 in the presence of a range of concentrations of human serum albumin. This protocol is based on that described by Giuliano et al. who measured intrinsic drug clearance in a microsomal stability assay (Giuliano et al., 2005).

RESULTS
Step 1: Development of the Rat PBK Model IV Administration
A comparison of the default simulated and measured kinetics profiles of a single dose of 0.5 mg/kg homosalate is shown in Figure 1A. This shows that the default parameters resulted in a lower Cmax and AUC(0–24h) compared to the measured values in rats. Therefore, the automated parameter identification function in PK-Sim was used to optimize several homosalate-specific parameters to fit the observed concentration-time profile of homosalate in rats. These included, Fu and permeability between compartments, which were shown to significantly impact the distribution of homosalate and were therefore
optimized using predicted values based on compound properties (e.g., lipophilicity and molecular weight). The optimized Fu was determined to be 1.4%. The optimized permeability values were: interstitial-to-intracellular and intracellular-to-interstitial permeability (4.83 cm/min); endothelial permeability (1.9 cm/min); and blood cell-to-plasma partition coefficient (21.28). These optimized values resulted in a concentration-time profile of homosalate that was similar to that measured in rats after IV administration of 0.5 mg/kg (Figure 1B), with similar mean C<sub>max</sub> (320 ng/ml predicted compared to the measured value of 338.3 ± 106.8 ng/ml) and AUC<sub>(0-∞)</sub> values (123.1 ng h/ml predicted compared to the measured value of 113.6 ± 22.9 ng h/ml).

A sensitivity analysis was conducted using AUC and C<sub>max</sub>. The outcome indicated that C<sub>max</sub> was more sensitive to several parameters than AUC (data not shown); therefore, a focus was made on C<sub>max</sub> to cover remaining influencing parameters. The results of a sensitivity analysis of the homosalate and physiological input parameters impacting the change in C<sub>max</sub> is shown in Figure 2A. This indicated that the C<sub>max</sub> is sensitive to several homosalate-specific parameters. The greatest relative change was observed for the Fu, with relative change of −0.77. As with the Fu, four other parameters (lipophilicity, plasma to blood permeability and blood/plasma partition coefficient and pKa) also negatively impacted the C<sub>max</sub>. By contrast, the blood/plasma permeability positively impacted the C<sub>max</sub> (with a ratio of 0.2). These influencing parameters altered the C<sub>max</sub> within one SD. Consequently, the estimated kinetics parameter uncertainty was set at the 5th to 95th C<sub>max</sub> around the mean value, where the SD around the mean defined the range of the influencing parameters.

The mean and CI 95% of AUC<sub>(0-∞)</sub> and C<sub>max</sub> of the additional doses, 2 and 5 mg/kg, were also estimated using the optimized model and were shown to be similar to in vivo values (shown in Figure 3, along with the comparison of the dose of 0.5 mg/kg). The simulated kinetics indicated a linear correlation between the

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**FIGURE 1** Predicted versus observed (blue circles) concentration-time profile of homosalate after an IV dose of 0.5 mg/kg. Profiles were generated using (A) default values and (B) optimized values. The blue circles represent measured values and the red line denoted a simulated profile.
AUC_{0-\infty}\) and \(C_{\text{max}}\) with the dose, which was in accordance with \textit{in vivo} findings. These results showed that the model performed well and could be used to extrapolate to the oral route.

**Oral Administration**

The optimized PBK rat model was used to estimate the \(C_{\text{max}}\) and \(\text{AUC}_{(0-24)}\) of homosalate after single and repeated oral administration of 60, 120, 300 and 750 mg/kg/day to rats. This would enable an understanding of the internal concentration across this dose range and; therefore, provide a more relevant representation of exposure than the applied dose (in mg/kg) to compare with human exposure (SCCS, 2021). The model-based approach used by Thelen et al. (Thelen et al., 2011) to calculate the intestinal permeability coefficients predicted the oral absorption (\(F_a\)) to be almost complete (i.e., 100%); however, the oral bioavailability (\(F_b\)) was predicted to be 83%, which might be due to a first-pass effect. However, the SCCS recommends a value 50% of the administered dose for chemicals without measured kinetics; therefore, this value was used as a more conservative approach which could be considered to incorporate a higher first-pass effect (since a lower systemic concentration would be linked to any adverse effects). The
intestinal permeability was adjusted to result in an oral bioavailability (Fb) of 50%. As a result, the corresponding oral absorption (Fa) was estimated at 81%. Due to the lack of available data, absorption kinetics was kept as default; however, both internal exposure metrics (short term: Cmax and long term: AUC) were reported and can be implemented for the calculation in the safety assessment. The predicted Cmax and AUC(0–24 h) values for each of the doses are shown in Table 3.

### Step 2: Development of the Human PBK Dermal Model

A preliminary sensitivity analysis was conducted on the skin penetration model (Dancik et al., 2013) to assess the effect of varying the main model parameters on the cumulative permeant mass (Q) that crosses partially hydrated skin over a 24 h period. The parameters tested were the partition coefficients with respect to water and the diffusion coefficients of each skin layer. The physicochemical properties specific to the permeant (homosalate) were used as input into the model, which, as a first step, calculates the nominal partition and diffusion coefficients of the compound. The stratum corneum was identified as the main barrier to homosalate skin penetration, with a negligible contribution of the epidermis and dermis to the overall cumulative mass penetration. Thus, the initial sensitivity analysis demonstrated that the cumulative mass, Q, of homosalate crossing the skin was most sensitive to the properties of the stratum corneum. The relevant system-dependent parameters were thus identified as those that comprise the stratum corneum partition coefficient with respect to water (Ksc/w) and the stratum corneum diffusion coefficient (Dsc). The derivation of these parameters from the parameters in the model are discussed by Danick et al. (Dancik et al., 2013). The variability in the skin model parameters translates to a significant variation in Q, which can impact the whole-body permeant disposition. When default input values were used, the skin absorption was predicted to be 0.6% of the applied dose, which is 3-fold lower than that was considered for a risk assessment (SCCP, 2007). Thus, three pharmacokinetic analyses were therefore conducted: In the first, the parameters of the skin permeation model were set to generate a cumulative mass penetration equal to Q = 2.07 µg/cm²/24 h, representing 2% skin penetration of homosalate. In the second and third analysis, the parameters were set to deliver 4 and 5.46 µg/cm²/24 h, representing 3.86 and 5.3% of applied homosalate, respectively. The cumulative amount of homosalate (Q) entering the bloodstream per cm² over 24 h, excluding first-pass skin metabolism, was incorporated using these values, and ranged from 2.07 to 5.46 µg/cm² (Table 4). The corresponding Cmax and AUC(0–24 h) values were then estimated by the human whole-body PBK model, which ranged from the mean of 2.38–6.3 ng/ml and 49.39–130.4 ng h/ml, respectively. The linear relationship between the Cmax and the dose, reported in rats, was also observed in the human dermal PBK model.

The results of a sensitivity analysis of the impact of homosalate-specific and physiological input parameters on the change in Cmax in human plasma after topical application is shown in Figure 2B. This indicated that, apart from the organism-specific input parameters, the Cmax was sensitive to several homosalate-specific parameters. As with the rat PBK model, the Fu negatively impacted the Cmax (with a ratio of ~0.38), although the greatest relative negative change was observed for the lipophilicity, with relative change of ~1.8. Three other parameters (plasma/blood cell permeability, blood cells/plasma partition coefficient and pKa) also negatively impacted the Cmax. As for the rat PBK model, the blood/plasma permeability increased the Cmax (with a ratio of 0.55). The same set of homosalate parameters was therefore used to calculate the uncertainty of Cmax following dermal exposure (based on the SD of the influencing input parameters). The resulting developed models were applicable for a range of dermal penetration of homosalate from 2 to 5.3%, including uncertainty resulting from formulation effects.

### Table 3: Predicted kinetics parameters of homosalate in rats after repeated oral administration.

| Dose (mg/kg/day) | Cmax (µg/ml) CI95% | AUC(0–24 h) (µg h/ml) CI95% |
|-----------------|-------------------|-------------------------------|
| 60              | 932.4 (769.21–1192) | 16174.05 (13295.64–20953.58) |
| 120             | 1837.8 (1512–2339)  | 31979.89 (26443.22–40847.898) |
| 300             | 4905 (4084–6188.8) | 84295 (70787–104639) |
| 750             | 11543 (9581–14545) | 200744 (167901–250263) |

### Table 4: Impact of the dermal delivery value on the estimated Cmax of homosalate after a single dermal administration. For each scenario, a single dose of 10% homosalate was applied to the whole-body surface (17500 cm²) at 103 µg/cm²/ day. The values for the Day 1 kinetics.

| Dermal delivery (% of the applied dose) | Cmax (µg/ml) mean | AUC(0–24 h) (µg h/ml) mean |
|----------------------------------------|-------------------|--------------------------|
| 2%                                     | 2.07              | 49.39                     |
| 3.86%                                  | 2.38              | 49.39                     |
| 5.3%                                   | 4.13              | 130.40                    |

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The human PBK dermal model performance was evaluated by comparing predicted values with measured values over 24 h from a clinical trial (Matta et al., 2020). When the dermal penetration was set at the lowest value of 2% of the applied dose, the mean C\text{max} (3.72 ng/ml) was comparable to that observed in humans (4.6 ng/ml). However, the simulated C\text{max} CI95% (6.45 ng/ml) accounted for only \(\sim 73.3\%\) of the clinical C\text{max} values (which ranged from 2 to 8.8 ng/ml). Therefore, the dermal penetration value was adjusted to result in a more comparable C\text{max} (mean = 4.55, CI95% = 7.88 ng/ml), which covered 89.6% of the clinical values (Figure 4A). The corresponding dermal penetration was estimated to be 2.48% of the applied dose. The plasma T\text{max} of 10–12 h observed in humans was also predicted by the dermal model.

**Step 3: Simulation of Kinetics Following Administration to Human Virtual Populations**

The optimized human PBK dermal model was used to estimate C\text{max} and AUC\text{(0–24)} values after repeated bi-daily dermal exposure of product containing 10% homosalate for 30 days. The simulated concentration-time profile of homosalate following repeated dermal exposure over 30 days, for which
the dermal penetration was estimated to be 5.3% of the applied dose, as shown in Figure 4B. The plasma concentration was predicted to rise rapidly in the first 24 h and then slowly increase after this time. Table 5 shows the impact of using different values for dermal penetration on the internal exposure to homosalate present at 10% in a product. The mean Cmax ranged from 5.22 ng/ml using 2.0% of the applied dose to 13.62 ng/ml when using 5.3% of the applied dose. The mean AUC(0–24) ranged from 119.81 ng h/ml using 2.0% of the applied dose to 316.49 ng h/ml when using 5.3% of the applied dose.

DISCUSSION

In common with many jurisdictions, the European cosmetics industry fully supports the animal testing ban under the EU Cosmetics Products Regulation. The industry has decades of commitment in promoting the use of alternatives to animal testing for safety assessment of cosmetics and has been at the forefront of the development and use of non-animal testing methods for safety assessment. One approach to allow animal-free decision-making is “Next generation risk assessment” (NGRA), which is an exposure-led and hypothesis-driven tiered workflow designed to prevent harm to humans (Dent et al., 2021). One of the principles underpinning the NGRA of cosmetic ingredients is that it is only conducted following an appraisal of all existing information, which may already be sufficient to make a safety decision. This is an important principle because many cosmetic ingredients have been used safely for decades and it would not be sensible to ignore historical animal data generated prior to the animal testing ban. This is the case for homosalate, which was the subject of historical animal tests which need to be put into context for the overall safety evaluation. One of the critical tools that allows this is PBK modelling. Here, PBK models were built and refined to estimate homosalate plasma concentrations over time, first, in rats after repeated oral exposure to a dose related to a no effect level observed in a toxicity study and, second, in humans following consumer-relevant dermal exposure as a sunscreen to the whole-body. The plasma values (e.g., mean, CI5% or CI95%, depending on the level of conservatism and the interpretation of the safety assessor) can then be compared to calculate a MoIE, which should ideally be above a safety factor of 25 to be considered protective of human health. This safety factor is based on the safety factors described by WHO (World Health Organization, 2005), which account for an overall uncertainty of 25 due to animal to human differences in toxicodynamics (safety factor = 2.5), human variability in toxicodynamics (safety factor = 3.16) and human variability in toxicokinetics (safety factor = 3.16). An overview of the development of the PBK models is shown in Figure 5, along with their associated assessments for model qualification and the derivation of the MoIE.

The PBK models were developed in a stepwise manner. The first step was to build a PBK rat model, which was optimized to simulate the kinetics of a single IV dose of 0.5 mg/kg homosalate administered to rats. While it is understood that the use of animal data is not currently possible in the cosmetics industry, legacy data generated prior to March 2013 can be used if available. Since the experimental work of the rat study reported by Kim et al. was conducted prior to the animal testing ban, it could be used for the purpose of model building and qualification. In the absence of the in vivo data, our study showed that the default values of the PK-Sim model would underpredict the internal exposure (AUC(0–24 h) and Cmax) to homosalate. This would be considered a conservative model, since toxicity observed would be linked to a lower plasma concentration that was actually present. The optimization using automated parameter identification function in PK-Sim ensured that a more realistic prediction of plasma concentrations was obtained. The accuracy of the refined model was confirmed by comparing simulated and measured kinetics profiles of two additional IV doses in rats.

The IV model was then adapted to incorporate intestinal permeability to enable the extrapolation of the NOAEL in a repeated oral dose study in rats to a plasma concentration. Two different values for bioavailability were considered but since the value of 50% recommended by the SCCS leads to a lower plasma concentration than the in silico derived oral bioavailability value of 83%, the former was incorporated to consider biologically plausible first-pass metabolism and to ensure a more conservative value. This was also considered prudent because measured data were unavailable for comparison with the resulting kinetics profile.

A PBK human dermal model was then developed to estimate blood concentrations following repeated topical exposure of humans to a product containing 10% homosalate. This value can be compared with the plasma concentration at the NOAEL to calculate the MoIE for the risk assessment (using mean values of AUC(0–24 h), see Figure 4). This allows the determination of a safe dose of homosalate based on a systemic exposure-based risk assessment rather than an external exposure-based risk assessment.

### Table 5 | Estimated kinetics parameters of homosalate after repeated dermal exposure over 30 days and different dermal penetration potential. For each scenario, 10% homosalate was applied to the whole-body surface (17500 cm²) twice a day (18 g cream per day, 103 μg/cm²/day). The same set of the influencing homosalate-specific input parameters and the population modeling were implemented to calculate the Cmax uncertainty. The values for the Day 30 kinetics.

| Dermal penetration (% of the applied dose) | 2.0  | 2.5  | 3.86 | 5.3  |
|--------------------------------------------|------|------|------|------|
| Cmax (mean) (ng/ml)                        | 5.22 | 6.65 | 9.94 | 13.62|
| Cmax (CI5-95%) (ng/ml)                     | 3.16–8.17 | 4.02–10.4 | 6.01–15.53 | 8.24–21.27|
| AUC(0–24h) (mean) (ng.h/ml)                | 119.81 | 153.02 | 230.96 | 316.49|
| AUC(0–24h) (CI5-95%) (ng.h/ml)             | 72.6–187.09 | 92.72–238.96 | 139.92–360.68 | 191.73–494.26|
The refined dermal model was suitable for varying extents of dermal penetration, although the value that best reflected the plasma concentrations measured in humans was 2.5% of the applied dose. When considering a risk assessment, the highest value that considers formulation effects (i.e., 5.3% of the applied dose) can be considered to be the most conservative.

The WHO and OECD Test Guidelines require a PBK model to be valid and applicable to the purpose of its use. The model validity (i.e., reliability and relevance) is based on several factors, including (1) a biological basis of the model structure and parameters, (2) a theoretical basis of the model equations, (3) reliability and relevance of input parameters, and (4) the sensitivity of the output to input parameters (WHO, 2010; OECD, 2021). For the first two requirements, the PK-Sim is an ideal platform with a high level of confidence, since it is an open-source model for which the codes, equations parameters, structure and assumptions are freely available. It is also accepted by the FDA for waiving of clinical trials (Zhuang and Lu, 2016). Several assumptions were implemented in the PBK model e.g., it assumes perfusion rate/blood flow limited kinetics for homosalate within the organs and tissues, which describes the kinetics for small lipophilic molecules. The developed model consistently reproduced the general kinetics of the parent molecule under various conditions i.e., different doses, routes of exposure (IV, dermal exposure), ultimately simulating kinetics profiles for the intended applications in rats and humans.

An important requirement in building a PBK model to confirm its validity is to subject it to a sensitivity analysis to identify the parameters which markedly influence the predictions. A model is considered to be reliable when a small change in a parameter value leads to changes in predictions of a dose metric that are less than the variation expected from its experimental measurement (WHO, 2010; OECD, 2021). The PBK model here was shown to be sensitive to parameter changes, whereby sensitivity analysis identified several homosalate-specific parameters (calculated partition coefficient and permeability blood cells/plasma) which influenced the C_max within one SD of the mean. The uncertainty of the in vitro measured parameters (Fu and metabolic clearance) was considered within the SD. The skin penetration model was optimized to recapitulate in vitro data and, due to the variation of values of homosalate dermal penetration (2–5.3% of the applied dose, presumably due to formulation effects and the inherent variability between skin penetration studies using different donors), several skin models were built to address dermal penetration uncertainty. The uncertainty calculated in the current study covers both true uncertainty (homosalate-specific parameters) and variability (population variability). Uncertainties in the whole-body PBK model were considered via population modeling, where a virtual population was built to cover variations in the anatomy and physiology of humans and rats. This covered the variation of the organ volumes, organ composition, tissue and body fluid physiology and blood flow rates.

The degree of confidence in the predictions of dose metrics by a PBK model depends upon how well it has been tested against measured data from a variety of dosing scenarios (WHO, 2010). The level of confidence in the model described here was considered to be medium-to-high for rat kinetics and high for human kinetics, since it was able to simulate the general trend of the concentration-time profile (peaks and decline) of homosalate under different exposure scenarios in humans and rats. This was evident upon visual inspection and, quantitatively, with respect to the plasma concentrations. While this was possible in the current study, PBK
models for other compounds may not be possible and would rely on the quality of the in vitro data used as input, as well as the use of sensitivity analysis to assess the confidence for a specific purpose (OECD, 2021).

In conclusion, we have developed PBK models which are able to correctly predict the internal exposure of homosalate according to different exposure scenarios. There was a medium-to-high level of confidence in the model output according to the classification defined in guidance documents (WHO, 2010; OECD, 2021) and, therefore, is useful in risk assessment. The process established here may serve as a reference supporting future homosalate PBK studies (e.g., extrapolation across dose-routes, between species, from high to low dose levels, and over various dosing scenarios), in further safety assessments on homosalate exposure. Indeed, in the absence of in vivo data, such human PBK models will be the heart of future completely non-animal risk assessments; therefore, valid approaches, as used here, will be key in gaining their regulatory acceptance.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

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AUTHOR CONTRIBUTIONS

All authors contributed to conception and design of the PBK modeling. SG and LR generated ADME data used as input for the model. AN performed the PBK simulations. NH wrote the first draft of the manuscript. All authors contributed to manuscript revision, read, and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fphar.2021.802514/full?_supplementary-material
