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# How is a global sensitivity analysis of a catchment-scale, distributed pesticide transfer model performed? Application to the PESHMELBA model

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**Abstract.** Pesticide transfers in agricultural catchments are responsible for diffuse but major risks to water quality. Spatialized pesticide transfer models are useful tools to assess the impact of the structure of the landscape on water quality. Before considering using these tools in operational contexts, quantifying their uncertainties is a preliminary necessary step. In this study, we explored how global sensitivity analysis could be applied to the recent PESHMELBA pesticide transfer model to quantify uncertainties on transfer simulations. We set up a virtual catchment based on a real one, and we compared different approaches for sensitivity analysis that could handle the specificities of the model: a high number of input parameters and a limited size of sample due to computational cost and spatialized output. After a preliminary screening step, we calculated Sobol' indices obtained from polynomial chaos expansion, Hilbert–Schmidt independence criterion (HSIC) dependence measures and feature importance measures obtained from random forest surrogate model. Results from the different methods were compared regarding both the information they provide and their computational cost. Sensitivity indices were first computed for each landscape element (site sensitivity indices). Second, we proposed to aggregate them at the hillslope and the catchment scale in order to get a summary of the model sensitivity and a valuable insight into the model hydrodynamic behaviour. Conclusions about the advantages and disadvantages of each method may help modellers to conduct global sensitivity analysis on other such modular and distributed hydrological models as there has been a growing interest in these approaches in recent years.

## 1 Introduction

Pesticide transfers from fields to water bodies are a major but also complex environmental concern. Significant efforts are required to assess risks for aquatic ecosystems and human lives. To do so, numerical models that simulate pesticide transfers and fate are necessary tools to support risk management. Among others things, such models make it possible to explore and compare scenarios of exposure and to assess mitigation measures. For this purpose and to support decision-making, physically based models such as in Reichenberger et al. (2007) or Dosskey et al. (2011) are particularly valuable. When they are distributed at the scale of the catchment, these models also make it possible to take into account the landscape configuration. This is of particular interest as the landscape configuration is of big influence on transfers and regulation, and corrective actions can thus be relevantly set up at this scale. PESHMELBA (Rouzies et al., 2019) is such a modular, process-based and distributed model that simulates pesticide transfers and fate at the scale of small agricultural catchments and that takes into account the impact of the landscape composition. However, PESHMELBA is characterized by a complex structure, numerous couplings and interactions that raise additional challenges about diagnosing model behaviour and uncertainty quantification (Gupta et al., 2008; van Griensven et al., 2006). To address the issue of uncertainty quantification, sensitivity analysis is a powerful tool that is being increasingly used in environmental models (Hamby, 1994; Tang et al., 2007; Nossent et al., 2011; Garcia et al., 2019; Alipour et al., 2022). Among its objectives, sen-

sitivity analysis contributes to obtaining greater insight about the model behaviour. It also contributes to identifying which input factors should be best characterized so as to significantly reduce the total uncertainty.

In the field of pesticide transfer modelling, Dubus et al. (2003) and Holvoet et al. (2005) first performed one-at-a-time (OAT) sensitivity analysis, meaning that each parameter influence was scrutinized individually, while other parameters were set to their nominal values. Although simple and computationally cheap to set, results from OAT analysis may be inaccurate in the case of a non-linear input–output relationship (Saltelli et al., 2004; Nossent and Bauwens, 2012). As a result, global sensitivity analysis (GSA) that varies all input factors simultaneously, on their entire ranges of definition, is favoured to address the sensitivity of environmental models. A large range of GSA methods exist which are not theoretically nor practically equivalent to set, and the method should be carefully chosen depending on the application characteristics (Song et al., 2015; Pianosi et al., 2016; Sarrazin et al., 2016). In the field of pesticide transfer modelling, studies mainly use variance decomposition methods. This approach is particularly appreciated as it allows us to characterize not only the contribution of each input factor individually but also the contributions of interactive effects between input factors. In addition, sensitivity indices calculated from variance-based methods are easy to interpret because they represent the portion of output variance that can be apportioned to an input factor (Saltelli, 2002). The Sobol' method (Sobol, 1993) is the most popular variance-based method, and it has been widely used in pesticide studies (e.g. Hong and Purucker, 2018; Gatel et al., 2019; D'Andrea et al., 2020; Faúndez Urbina et al., 2020). In some other studies (Fox et al., 2010; Lauvernet and Muñoz-Carpena, 2018), variance decomposition is also performed based on Fourier transform methods (FAST and eFAST methods). However, such variance decomposition methods are characterized by a high computational cost (meaning that they require a lot of model evaluations to compute sensitivity indices) that cannot always be afforded. As the computational cost quickly increases with the number of input factors, a classical approach consists in first applying a *screening* step to identify (if there are) parameters that have a negligible influence on the output variability. Non-influential parameters are then removed or set to constant values. Once the input space dimension has been reduced, variance decomposition can be applied on the reduced set of input factors to classify remaining input factors according to their relative contribution to output variability (*ranking* step).

Table 1 summarizes the characteristics of the GSA application for the previously cited pesticide studies (such a table is meant to provide some typical examples rather than be an exhaustive list of studies on the field of pesticide modelling). It shows that in these studies, the input space scrutinized for ranking contains at most 24 parameters (in some cases after a screening step) and that the ranking step always

uses more than 10 000 simulations. However, in the case of the PESHMELBA model (and probably of other distributed, physically based, hydrological and pesticide models) a classical application involves around 150 parameters. In addition, 10 000 simulations may be very hard to reach in realistic applications due to computation time. Such characteristics make classical approaches for variance-based GSA quite hard to apply to PESHMELBA. Then, there is a need for exploring other, low-computational-cost GSA methods to perform a ranking task in the PESHMELBA model. Recently, new GSA approaches that require much less model evaluations have emerged. For instance, computing a metamodel based on polynomial chaos expansion on the model (Ghanem and Spanos, 1991) allows us to compute Sobol' sensitivity indices directly, for a very limited computational cost (Sudret, 2008; Fajraoui et al., 2011; Wang et al., 2015). From a totally different point of view, the Hilbert–Schmidt independence criterion (HSIC) proposed by Gretton et al. (2005b) is used as a sensitivity measure in Da Veiga (2015). It describes the global dependence between the output and each input factor from a probabilistic point of view, also from a very limited computational budget. Finally, the growing interest for machine learning techniques is paving the way for new approaches of GSA, such as the random forest method (RF). Its structure provides valuable information on feature importance that can be processed as sensitivity indices like in Harper et al. (2011) and Aulia et al. (2019) (see Antoniadis et al., 2021, for a review on the use of random forests for sensitivity analysis). However, such innovative methods have never been applied to a complex, distributed pesticide transfer model before.

Therefore, the objective of this paper is to evaluate and to compare three new, low-computational-cost GSA methods for application to the PESHMELBA model. The methods will be especially compared in terms of interpretability of the sensitivity indices they provide and reliability (based on the study of their convergence rate). We also investigate whether such approaches suit the spatialized aspect of PESHMELBA. To do so, we investigate the relevancy of computing both local and aggregated indices following the recommendations from Saint-Geours (2012). The analysis is performed on a virtual scenario based on a real catchment in the Beaujolais region (France). The paper is organized as follows: we describe the PESHMELBA model in Sect. 2.1 and the model setup in Sect. 2.2. The input sampling is described in Sect. 2.3, and then we introduce the different GSA approaches and the methodology used for landscape analysis from Sect. 2.4 to Sect. 2.6. Results are presented in Sect. 3. They focus successively on screening (Sect. 3.1), comparison of GSA methods (Sect. 3.2) and spatial analysis (Sect. 3.3). Finally, Sect. 4 gathers some points of discussion.

**Table 1.** Examples of global sensitivity analysis performed on pesticide transfer models.

Reference	Model name	No. of input factors for ranking	Method	No. of model evaluations
Fox et al. (2010)	VFSMOD	18	eFAST	14 977
Lauvernet and Muñoz-Carpena (2018)	VFSMOD	24	eFAST	75 544
Hong and Purucker (2018)	PRZM	11	Sobol'	195 000
Gatel et al. (2019)	CATHY	15	Sobol'/Sobol' with ROA-LHS sampling (Tissot and Prieur, 2015)	1922/17 000
D'Andrea et al. (2020)	PWC	24	Sobol'	11 776
Faúndez Urbina et al. (2020)	SWAP-PEARL	< 14	Sobol'	> 18 000

## 2 Material and methods

### 2.1 The PESHMELBA model

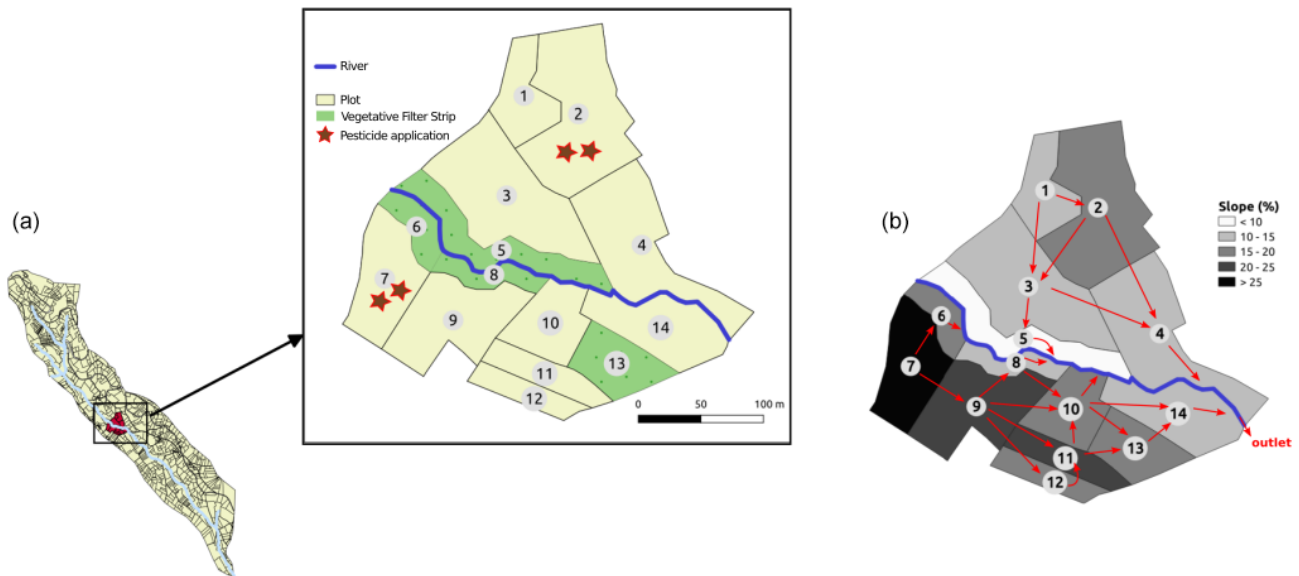
The PESHMELBA model represents a catchment as a set of interconnected components that stand for landscape elements such as plots, vegetative filter strips (VFSs), ditches, hedges or rivers (Rouzies et al., 2019). In order to respect the spatial organization and the heterogeneity of the landscape, it deals with mesh elements that can be surfaces or lines. Surface mesh elements are called homogeneous units (HUs). A HU is a portion of landscape that is homogeneous in terms of hydrodynamic processes and agricultural practices. Linear mesh elements are called reaches. A reach is characterized by its nature (so far ditch, river or hedge) and by its neighbouring components: it is at most in contact with one elementary mesh element on each bank. In addition to its geometric or hydrodynamic properties, each mesh element is characterized by its one-way connections with the neighbouring components that stand at a lower altitude. One or several processes are represented on each element depending on its nature. Lateral transfers at surface and in the subsurface between elements are also integrated. Independent codes called units are used to simulate the different processes, depending on the knowledge the user has on the targeted catchment functioning. Then, the OpenPALM coupler (Fouilloux and Piacentini, 1999; Buis et al., 2006) is used to couple the units and to build the complete application. OpenPALM has adapted features to easily deal with spatial and temporal aspects of the model. For example, synchronization tools make it possible to couple processes with different time steps. The final structure is highly modular, and process representations can easily be added, upgraded or removed depending on the landscape description. These features make PESHMELBA particularly suitable for scenario exploration. PESHMELBA focuses on surface and subsurface transfers of water and pesticides. An extensive description of elements and processes already included can be found in Rouzies et al. (2019). The PESHMELBA version used in this study integrates a representation of water and pesticide transfers on plots, VFSs

and rivers. Each plot or VFS is represented by a unique column of soil divided into vertical cells. In such a column, vertical infiltration is simulated using a solution of the 1D Richards equation proposed by Ross (2003). An adapted set of parameters makes it possible to represent high infiltration rate, surface runoff reduction, and enhanced adsorption and degradation on VFSs. Root-water uptake is integrated based on Varado et al. (2006). Surface runoff routing is represented based on the kinematic wave (Lighthill and Whitham, 1955), and the Darcy law (Darcy, 1857) is used for lateral subsurface transfers. In addition to shallow groundwater tables, PESHMELBA also represents shallowly perched water tables and associated lateral transfers. Finally, the reactive transfer of solutes is represented: advection and degradation, based on a first-order law, and adsorption, based on linear or Freundlich isotherms, are integrated. Each river or ditch reach is represented by a unique tank. The River1D module (Branger et al., 2010) solves the kinematic wave equation for water routing and pesticide advection in the network. Groundwater–river exchanges are represented by the Miles formulation adapted by Dehotin et al. (2008).

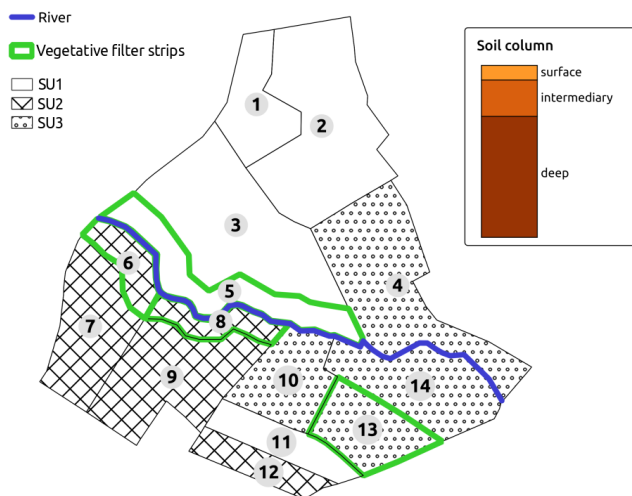
### 2.2 Model setup

A virtual scenario of limited size is set from a portion of La Morcille catchment (France) in order to explore the different GSA methods and to ease interpretation of spatialized results. The chosen portion is selected so as to remain representative of the global composition of La Morcille catchment in terms of soil, slope, type and size of elements, as well as interface length between them. The chosen scenario is composed of 10 vineyard plots, four vegetative filter strips and five river reaches that delimit a left and a right slope (see Fig. 1).

Soils types on the catchment are mainly sandy (Peyrard et al., 2016). We use the classification from Frésard (2010) that groups soil types into three main soil units (SUs). Each SU is defined by the vertical succession of three or four soil layers, also called soil horizons: one surface horizon, one or two intermediary horizons, and one deep horizon, as depicted



**Figure 1.** (a) Portion of La Morcille catchment selected to perform sensitivity analysis. Yellow units stand for vineyard plots, while green units stand for vegetative filter strips. Brown stars denote locations of pesticide application. (b) Slopes and connections between elements.



**Figure 2.** Soil type locations for the case study. Green contours show the vegetative filter strips.

in Fig. 2. Note that interface depths can vary from one SU to another. The reader may refer to Rouzies et al. (2019) for further details on how soil types and soil horizons are represented in PESHMELBA. At the catchment scale, the classification results in the following SU (see Fig. 2): sandy soil (SU1), sandy soil on clay on the right bank plateau (SU2), and heterogeneous sandy soils on lower slopes and thalwegs (SU3). Spatial arrangement is set in order to be as realistic as possible in terms of possible interfaces between SUs. Each SU is set at least on one vineyard plot and one VFS on the virtual scenario.

Considering the different soil horizons whose hydrodynamic behaviour and texture must be parameterized, the two types of vegetation (grassland and vineyard), and the different landscape element types (plots, VFSs and river reaches) that are simulated, the scenario results in 145 input parameters to be considered for sensitivity analysis. They are described in Table 2 as well as the spatial level on which they are set and their values for the nominal simulation.

For the nominal scenario, values for bulk density  $bd$  and organic carbon content  $moc$  are available from Van den Bogaert (2011) and Randriambolohasinirina (2012). They are, as well as hydrodynamic parameters for each soil type, described in Table 3. Retention values measured by Van den Bogaert (2011) are used to fit retention curve using the SWR-Cfit tool (Seki, 2007). A Schaap–van Genuchten conductivity curve is used (Schaap and van Genuchten, 2006; Ross, 2006), whose matching points at saturation  $K_0$  and empirical pore connectivity  $L$  are derived from conductivity data and retention parameters from retention curve fitting. Surface organic carbon content is set equal to that of the first soil horizon on plots and VFSs. For each SU, only the first soil horizon on VFSs differs from vineyard plots so as to highlight enhanced infiltration capacities. The surface horizon on VFSs is characterized by a 2.8 % organic carbon content (Randriambolohasinirina, 2012) and a saturated hydraulic conductivity of  $150 \text{ mm h}^{-1}$  (or  $4.31 \times 10^{-5} \text{ m s}^{-1}$ ) following Catalogne et al. (2018). In the absence of data to characterize potential anisotropy of vertical and horizontal saturated conductivities  $K_{sv}$  and  $K_{sh}$ , isotropy is considered; thus the ratio  $K_{sh}/K_{sv}$  is set to 1 in the catchment.

The pesticide chosen in this study is tebuconazole as it is a fungicide widely used in the La Morcille catchment.

**Table 2.** Input factor description, corresponding spatial level definition and value for the nominal scenario. Nominal values are explicitly distinguished between vineyard plots and VFSs with the character “–” when needed.

Input factor [units]	Description	Spatial level definition	Nominal value
Soil parameters			
thetas [ $\text{m}^3 \text{m}^{-3}$ ]	Saturated water content	Soil horizon	see Table 3
thetar [ $\text{m}^3 \text{m}^{-3}$ ]	Residual water content	Soil horizon	see Table 3
Ks [ $\text{m s}^{-1}$ ]	Saturated hydraulic conductivity	Soil horizon	
hg [m]	Air-entry pressure in van Genuchten retention characteristic curve	Soil horizon	see Table 3
mn	Deduced parameter from van Genuchten retention characteristic curve $n$ : $mn = n - 1$	Soil horizon	see Table 3
Ko [ $\text{m s}^{-1}$ ]	Matching point at saturation in modified Mualem–van Genuchten conductivity curve (Schaap and van Genuchten, 2006)	Soil horizon	see Table 3
$L$	Empirical pore-connectivity parameter	Soil horizon	see Table 3
bd [ $\text{g cm}^{-3}$ ]	Bulk density	Soil horizon	see Table 3
Pesticide parameters			
Koc [ $\text{m Lg}^{-1}$ ]	Freundlich sorption coefficient	Pesticide type	769
DT50 [d]	Half-life	Pesticide type	47.1
Vegetation parameters			
manning [ $\text{sm}^{-1/3}$ ]	Manning’s roughness	Vineyard plot–VFS	0.033–0.2
Zr [m]	Rooting depth	Vineyard plot–VFS	2.62–0.9
$F_{10}$	Fraction of the root length density in the top 10 % of the root zone	Vineyard plot–VFS	0.370–0.335
LAImin	Min LAI value	Vineyard (plot)	0.01
LAImax	Max LAI value	Vineyard (plot)	2.5
LAIharv	LAI value at harvest time	Vineyard (plot)	0.01
LAI	Constant LAI value	Grassland (VFS)	5
River parameters			
hpond [m]	Ponding height in the river bed	River reach	0.01
di [m]	Distance between the river bed and the limit of impervious saturated zone	River reach	1.5
Ks [ $\text{m s}^{-1}$ ]	Saturated conductivity of the river bed	River reach	$2.38 \times 10^{-5}$
manning [ $\text{sm}^{-1/3}$ ]	River bed Manning’s roughness	River reach	0.079
Plot and VFS parameters			
hpond [m]	Ponding height	Plot–VFS	0.01–0.05
adsorpthick [m]	Mixing layer thickness	All landscape elements (except river)	0.01

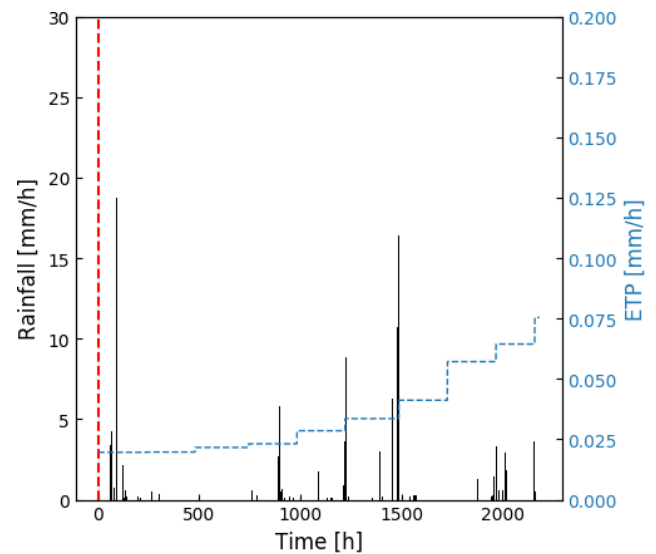
**Table 3.** Soil parameters for SU1, 2 and 3. Hydrodynamic parameters are based on the van Genuchten retention curve and on Schaap–van Genuchten conductivity curve fitting. Parameters are described in Table 2. Values for the surface horizon are explicitly distinguished between plot and VFS when they are different. Horizons 11, 12 and 13 are surface horizons for plots, whereas horizons 14, 15 and 16 are surface horizons for VFSs.

	Horizon	Depth [m]	Thetas [ $\text{m}^3 \text{m}^{-3}$ ]	Thetar [ $\text{m}^3 \text{m}^{-3}$ ]	hg [m]	$n$ [-]	Ks [ $\text{m s}^{-1}$ ]	Ko [ $\text{m s}^{-1}$ ]	$L$ [-]	bd [ $\text{g cm}^{-3}$ ]	moc [%]	
SU1	11–14	0.05	0.34	0.04	$-9.69 \times 10^{-2}$	1.27	$3.93 \times 10^{-5}$	$-4.31 \times 10^{-5}$	$2.86 \times 10^{-7}$	-8.43	1.34	0.91–2.80
	2	0.5	0.34	0.05	$-3.29 \times 10^{-2}$	1.20	$8.64 \times 10^{-5}$		$2.28 \times 10^{-7}$	-6.52	1.47	0.39
	3	0.65	0.32	0.08	$-2.09 \times 10^{-2}$	1.20	$5.39 \times 10^{-5}$		$7.47 \times 10^{-7}$	-4.24	1.57	0.10
	4	4	0.28	0.07	$-5.99 \times 10^{-2}$	1.23	$3.11 \times 10^{-5}$		$1.47 \times 10^{-6}$	-0.14	1.53	0.07
SU2	12–15	0.035	0.34	0.04	$-9.69 \times 10^{-2}$	1.27	$3.93 \times 10^{-5}$	$-4.31 \times 10^{-5}$	$2.86 \times 10^{-7}$	-8.43	1.34	1.15–2.80
	6	0.4	0.35	0	$-6.60 \times 10^{-2}$	1.13	$2.16 \times 10^{-5}$		$3.19 \times 10^{-7}$	9.66	1.59	0.68
	7	0.55	0.32	0	$-7.18 \times 10^{-2}$	1.08	$9.60 \times 10^{-6}$		$1.67 \times 10^{-7}$	-10	1.66	0.35
	8	4	0.42	0	$-3.02 \times 10^{-1}$	1.08	$3.98 \times 10^{-6}$		$9.72 \times 10^{-8}$	10	1.54	0.28
SU3	13–16	0.06	0.34	0.04	$-9.69 \times 10^{-2}$	1.27	$3.93 \times 10^{-5}$	$-4.31 \times 10^{-5}$	$2.86 \times 10^{-7}$	-8.43	1.34	0.75–2.80
	9	0.45	0.33	0.08	$-6.72 \times 10^{-2}$	1.26	$3.05 \times 10^{-5}$		$3.36 \times 10^{-7}$	0.42	1.46	0.37
	10	4	0.32	0.06	$-3.56 \times 10^{-2}$	1.18	$2.38 \times 10^{-5}$		$3 \times 10^{-7}$	1.05	1.62	0.40

It is a slightly mobile molecule, and we use a Freundlich isotherm to describe its adsorption equilibrium. Adsorption parameters are obtained from Lewis et al. (2016) ( $K_{oc} = 769 \text{ mL g}^{-1}$ , Freundlich isotherm exponent = 0.84). The surface degradation coefficient is also taken from Lewis et al. (2016) ( $DT_{50} = 47.1 \text{ d}$ ), and a decreasing degradation rate as a function of depth is set as in FOCUS (2001). A 500 g application is considered at the beginning of the simulation on plots 2 and 7 (see Fig. 1). Most of the transformation and adsorption of tebuconazole are supposed to happen on plots and VFSs at this modelling scale. Therefore, no adsorption or degradation is simulated in the river.

A no-flux boundary condition is applied on all sides except on the surface where rain and potential evapotranspiration are considered. Rain data are extracted from the BDOH database (Gouy et al., 2015). A 3-month simulation is performed for a time period characterized by long and intense rain events (670 mm cumulated), allowing for significant water and pesticide transfers both by surface runoff and subsurface saturated transfers. Potential evapotranspiration (PET) data are obtained from MétéoFrance for the neighbouring site of Liergues (MétéoFrance, 2008). Data are averaged over 10 d periods and corrected by a factor of  $-11\%$  to match the La Morcille site as recommended in Durand (2014) and Caisson (2019). Rain and PET data for the simulation are summarized in Fig. 3.

Although virtual, we aim at setting initial conditions as plausible as possible for this scenario. As running the model on a warm-up period is not possible due to data availability and computational cost limitation, initial water table levels are deduced from piezometric data on a neighbouring hillslope, and all soil columns are supposed to be in hydrostatic equilibrium at the beginning of the simulation. Data from several piezometers are available on a transect, perpendicular to the river. Data are extrapolated over the virtual hillslope



**Figure 3.** Climatic forcing (rain and potential evapotranspiration) for the simulation. The dotted red line stands for the one-shot pesticide application.

width on both sides of the river. An upstream  $0.177 \text{ m}^3 \text{ s}^{-1}$  flow is considered in the river based on local measurements (Gouy et al., 2015).

Two types of vegetation are represented in this scenario. Vineyard cover is considered on plots, while permanent grassland is simulated on VFSs. Considering the period of simulation (3 months), a fixed root depth ( $Z_r = 2.62 \text{ m}$ ) and a fixed root density in the first 10% of the root depth ( $F_{10} = 37\%$ ) are considered for vineyards following values reported in Smart et al. (2006) and confirmed by expert knowledge in the area. The root depth ( $Z_r$ ) is set to 0.9 m, and the root density in the first 10% of the root depth ( $F_{10}$ ) is set to 33.5% for grassland (Brown et al., 2007). For vineyards, the

leaf area index (LAI) is assumed to increase from a minimum value LAI<sub>min</sub> from leaves formation until a maximum value LAI<sub>max</sub> before declining until harvest LAI<sub>harv</sub>. Dates and associated values for this development cycle are taken from Brown et al. (2007), and they are detailed in Appendix A1. On grassland, the LAI is assumed to remain constant, and a nominal value of 5 is chosen based on Brown et al. (2007). In Table 2, the reader should note that the LAI parameter relates to the constant LAI value for grassland, while LAI<sub>min</sub>, LAI<sub>max</sub> and LAI<sub>harv</sub> relate to vineyard plots. The remaining parameters for the root extraction model are also fixed to nominal values proposed by Varado et al. (2006) and Li et al. (2001). Manning coefficients are set from data reported in Arcecent and Schneider (1989). A mature row crop value ( $0.033 \text{ s m}^{-1/3}$ ) is chosen for vineyards, while a high grass pasture value ( $0.2 \text{ s m}^{-1/3}$ ) is set for VFS cover.

Finally, ponding height is set to 0.01 m for vineyard plots, while an increased value is set for VFSs (0.05 m). According to Gao et al. (2004) and Walter et al. (2007), the surface mixing layer thickness is set to 0.01 m in both plot and VFS domains. In the river, the distance between the river bed and the limit of impervious saturated zone ( $d_i$ ) is set to 1.5 m (electrical resistivity tomography field measure, Rémi Dubois, personal communication, 2020), and the saturated hydraulic conductivity ( $K_s$ ) is set to  $2.38 \times 10^{-5} \text{ m s}^{-1}$  accordingly to local saturated conductivities in the neighbouring soil. The ponding height is set to 0.01 m, while the Manning coefficient is chosen equal to  $0.079 \text{ s m}^{-1/3}$ , as suggested in Arcecent and Schneider (1989) for channels with limited obstruction.

### 2.3 Input sampling

The input factor distributions are set to be as representative as possible of the available data on the catchment and the associated uncertainties. Mean values are taken from the nominal scenario described in Sect. 2.2. Distributions and standard deviations are assigned based on experimental measurements from the catchment of application, available scientific literature or expert knowledge. All assigned distributions and corresponding statistics are summarized in Appendix B.

As commonly found in the literature (Coutadeur et al., 2002; Fox et al., 2010; Schwen et al., 2011; Dairon, 2015; Dubus et al., 2003; Dubus and Brown, 2002), a lognormal distribution is assigned to the saturated hydraulic conductivity  $K_s$ . A 20 % coefficient of variation (CV) is used so as to remain representative of each soil horizon hydrodynamic behaviour. Distributions for Schaap–van Genuchten parameters could not be found in the literature; thus the empirical pore-connectivity parameter  $L$  is assigned a uniform distribution  $\pm 20\%$  around the mean value (Zajac, 2010). As the matching point at saturation in the modified Mualem–van Genuchten conductivity curve  $K_0$  has the same physical meaning as  $K_s$ , a lognormal distribution is also assigned to this parameter and a 20 % CV is set. Saturated water content

thetas, residual water content  $\theta_{\text{etar}}$ , van Genuchten parameter  $m_n$  and air-entry pressure  $h_g$  are assigned normal distributions (Schwen et al., 2011; Alletto et al., 2015; Dairon, 2015; Gatel et al., 2019). A 10 % CV is set for thetas (Gatel et al., 2019; Lauvernet and Muñoz-Carpena, 2018), and  $\theta_{\text{etar}}$  is assigned a 25 % CV (Gatel et al., 2019). A 10 % CV is set for  $m_n$  and  $h_g$  (Schwen et al., 2011; Alletto et al., 2015; Gatel et al., 2019). A uniform distribution is assigned to organic carbon content  $\text{moc}$  (Lauvernet and Muñoz-Carpena, 2018). A triangular distribution is assigned to the Freundlich sorption coefficient  $K_{oc}$  (Lauvernet and Muñoz-Carpena, 2018), and a normal distribution is assigned to the half-life  $DT_{50}$ . A 60 % CV is assigned to  $K_{oc}$  and  $DT_{50}$  distributions as such parameters are considered relatively uncertain (Dubus et al., 2003). Triangular distributions are assigned to Manning's roughness ( $manning$ ) on plots and for the river bed (Lauvernet and Muñoz-Carpena, 2018; Gatel et al., 2019). A uniform distribution with a  $\pm 20\%$  range around the mean value is assigned to remaining input factors as little information could be found in the literature (Zajac, 2010).

Using a fully distributed model such as PESHMELBA raises the issue of sampling strategy. Indeed, in this case study, even if the site is only composed of 14 surface units, the large number of soil horizons on the catchment, considering the hydrodynamic distinction between plots and VFSs, also dramatically increases the number of parameters. Sampling all parameters on each spatial unit leads to a huge number of simulations that could not be computationally afforded. Moreover, such independent sampling on a very large number of parameters may lead to misinterpretation of the sensitivity analysis results as the influence of physical processes could not be distinguished from spatial arrangement. For each sample, one set of soil parameters is therefore sampled for each soil horizon, and those parameters are applied to all spatial units that contain this horizon which sets the number of parameters to be considered in the GSA at 145.

### 2.4 Methodology for global sensitivity analysis in the PESHMELBA model

Although the PESHMELBA model is dynamic, model outputs considered in this paper are scalar quantities rather than temporal series to keep the problem simple. In order to investigate PESHMELBA abilities to properly represent transfers in a heterogeneous landscape, sensitivity analysis is performed on four hydrological and quality variables: (1) cumulated water volume transferred in the subsurface (saturated lateral transfers), (2) pesticide mass transferred in the subsurface (saturated lateral transfers) (3) cumulated water volume transferred on surface (surface runoff) and (4) cumulated pesticide mass transferred on surface (surface runoff). However, these quantities are spatialized, leading to multidimensional outputs. To deal with the spatialized aspect, GSA is first performed on scalar quantities, on each landscape element (see Sect. 2.4.1 to 2.4.3), then sensitivity indices are



aggregated in a second time to provide catchment-scale sensitivity indices (see Sect. 2.6).

The full workflow used to perform GSA in the PESHMELBA model is summarized in Fig. 4. Considering the high number of input parameters, a screening step is first performed to decrease the dimension of the problem. Screening is performed with a statistical independence test based on the HSIC measure (see Sect. 2.4.2) on an initial 4000-point Latin hypercube sample (LHS; McKay et al., 1979). Second, a new 2000-point LHS obtained from the reduced set of input parameters is computed to perform ranking. Sensitivity indices are computed from the 2000-point sample based on (1) variance decomposition (Sect. 2.4.1), (2) the HSIC dependence measure (Sect. 2.4.2) and (3) the feature importance measure obtained from random forests (Sect. 2.4.3). These methods are all global and model-free, and all suit non-linear, non-monotonic models. In addition, they all belong to the category of “given-data” methods which means that the input sample they consider does not require a specific structure. This is of particular interest for models that are computationally costly as it allows us to take advantage of pre-existing simulations. That is why such methods have known a growing interest in recent years (Saltelli et al., 2021; Sheikholeslami et al., 2021). However, such methods may not be equally costly to set, and they define the notion of sensitivity in contrasted ways. We compare them regarding the information they provide, their accuracy and their robustness.

In what follows, we denote  $Y \in \mathbb{R}$ , a given scalar output from PESHMELBA.  $Y$  is function of a multivariate input random vector  $Y = \mathcal{M}(X)$ , where  $X = (X_1, \dots, X_M) \in \mathbb{R}^M$  contains the 145 input parameters considered in this case study and where  $\mathcal{M}$  is the PESHMELBA model.

### 2.4.1 Variance decomposition

Variance-based methods aim at determining how input factors contribute to the output variance (Faivre et al., 2013). One of the most popular variance-based method is the Sobol’ method (Sobol, 1993). The Sobol’ indices capture the direct impact of any input and also describe the impact of input parameters in interaction with others. It is based on the decomposition of the total variance of the output:

$$\text{Var}[Y] = \sum_{s=1}^M \sum_{i_1 < \dots < i_s} V_{i_1, \dots, i_s}, \tag{1}$$

where  $V_{i_1, \dots, i_s}$  indicates the portion of variance that can be attributed to interactions between input parameters  $X_i, i \in i_1, \dots, i_s$ . From the above, one can define Sobol’ indices as

$$S_{i_1, \dots, i_s} = \frac{V_{i_1, \dots, i_s}}{\text{Var}[Y]}. \tag{2}$$

By definition,  $0 \leq S_{i_1, \dots, i_s} \leq 1$ . In particular, first-order sensitivity indices  $S_i = \frac{V_i}{\text{Var}[Y]}$  only account for the main effects

of parameter  $X_i$ . They can be interpreted as the decrease in the total output variance that could be obtained when removing the uncertainty about  $X_i$  when setting  $X_i$  to a fixed value (Tarantola et al., 2002). These indices are usually calculated as a first step as they often account for a large portion of the variance (Faivre et al., 2013). Total sensitivity indices  $S_{T_i}$  evaluate the total effect of an input factor  $X_i$  on the output by taking into account its main effect  $S_i$  and all interaction terms that involve it:

$$S_{T_i} = \sum_{\mathcal{I}_i} S_{i_1, \dots, i_s}, \quad \mathcal{I}_i = \{(i_1, \dots, i_s) \mid \exists k, 1 \leq k \leq s, i_k = i\}. \tag{3}$$

The total sensitivity index  $S_{T_i}$  stands for the portion of total output variance that remains as long as  $X_i$  stays unknown (Tarantola et al., 2002).

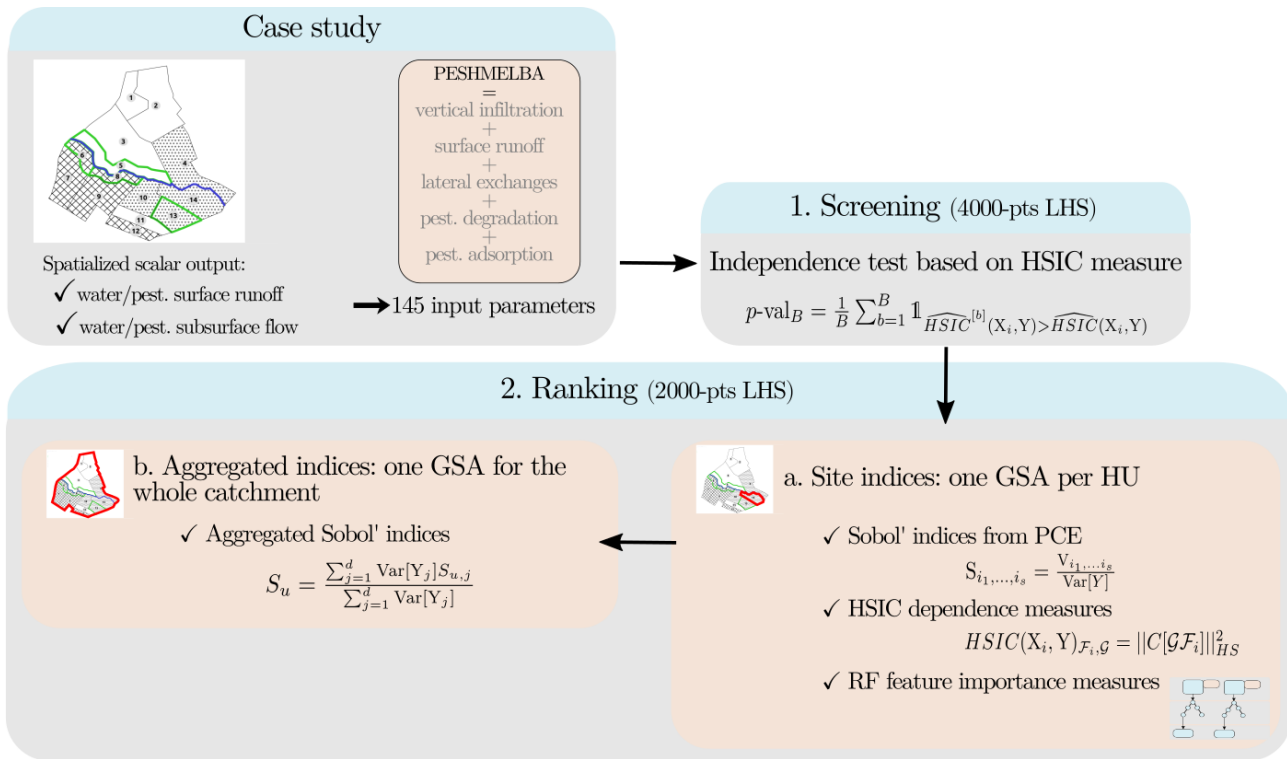
Sobol’ indices’ direct computation requires a large sample size that can not be afforded in this case study. As a result, we compute Sobol’ indices from a limited sample size, based on polynomial chaos expansion (PCE; Sudret, 2008) in order to circumvent such difficulty. This approach consists in building a surrogate model of which analytical polynomial expression is directly related to Sobol’ indices. Building a PCE and deducing the associated Sobol’ indices thus only requires a training sample of limited size and knowledge about each input parameter marginal distribution. More precisely, PCE provides a functional approximation of the computational model based on the projection of the model output on a suitable basis of stochastic polynomial functions in the random inputs (Ghanem and Spanos, 1991). For any square integrable scalar output random variable  $Y$ , its polynomial chaos expansion is expressed as follows:

$$Y = \sum_{\alpha \in \mathbb{N}^M} \gamma_\alpha \Psi_\alpha(X), \tag{4}$$

where the  $\Psi_\alpha$ ’s are multivariate orthonormal polynomials built according to the marginal probability density functions of each input factor, and  $\gamma_\alpha$  values are the associated coordinates. Expansion from Eq. (4) is usually truncated to a finite sum for practical computation, using for example a truncation scheme based on least-angle regression (Blatman and Sudret, 2011). The Sobol’ indices can then be obtained analytically from the coefficients  $\gamma_\alpha$  (see Sudret, 2008, for a demonstration of the relation between PCE and Sobol’ indices). In our study, the UQLab MATLAB software (Marelli and Sudret, 2014) is used to compute Sobol’ indices from the 2000-point LHS (step 2a, Fig. 4). We use a q-norm- and degree-adaptive sparse PCE based on the least-angle regression scheme (LARS; Blatman and Sudret, 2011), with q norm  $\in [0.1, 0.2, \dots, 1.0]$  and a maximum degree of 3.

### 2.4.2 The HSIC dependence measure

Sensitivity measures based on the Hilbert–Schmidt independence criterion (HSIC; Da Veiga, 2015) belong to the cat-



**Figure 4.** Full workflow used to perform GSA on the PESHMELBA model in three steps: (1) screening, (2a) ranking at a local scale and (2b) ranking at the catchment scale.

egory of dependence measures that quantify, from a probabilistic point of view, the dependence between each input and the output. The greater the dependency between the input factor and the output, the greater the associated sensitivity measure. The Hilbert–Schmidt independence criterion used for GSA is based on the cross-correlation between any non-linear transformations of some input factor  $X_i$  and the output  $Y$  (De Lozzo and Marrel, 2016). Such a dependence measure simultaneously captures a very broad spectrum of forms of dependency between the variables (Meynaoui et al., 2018). These indices can be estimated from small samples (a few hundreds of points) and do not depend on the number of inputs, which is a huge advantage.

The HSIC theory relies on reproducing kernel Hilbert space (RKHS) and kernel functions. Let  $\mathcal{F}_i$  denote the RKHS composed of all continuous bounded functions of input  $X_i$ , with values in  $\mathbb{R}$  and  $\mathcal{G}$  the RKHS composed of real-valued continuous bounded functions of output  $Y$  with values in  $\mathbb{R}$ .  $\langle \cdot, \cdot \rangle_{\mathcal{F}_i}$  (i.e.  $\langle \cdot, \cdot \rangle_{\mathcal{G}}$ ) is the inner product on  $\mathcal{F}_i$  (i.e.  $\mathcal{G}$ ), and  $k_{X_i}$  (i.e.  $k_Y$ ) is the corresponding kernel function that defines such a scalar product. The HSIC measure corresponds to the square of the Hilbert–Schmidt norm of the cross-covariance

operator  $C[\mathcal{GF}_i] : \mathcal{G} \rightarrow \mathcal{F}_i$ , which is

$$\begin{aligned} HSIC(X_i, Y)_{\mathcal{F}_i, \mathcal{G}} &= \|C[\mathcal{GF}_i]\|_{HS}^2 \\ &= \sum_{j,k} \langle u_j^i, C[\mathcal{GF}_i](v_k) \rangle_{\mathcal{F}_i} \\ &= \sum_{j,k} \text{cov}(u_j^i(X_i), v_k(Y)), \end{aligned} \quad (5)$$

where  $(u_j^i)_{j \geq 0}$  and  $(v_k)_{k \geq 0}$  are orthogonal bases of  $\mathcal{F}_i$  and  $\mathcal{G}$  respectively.

The resulting sensitivity indexes proposed by Da Veiga (2015) are defined for each input factor  $X_i, i \in \{1, \dots, M\}$  as

$$S_{X_i}^2 = \frac{HSIC(X_i, Y)_{\mathcal{F}_i, \mathcal{G}}}{\sqrt{HSIC(X_i, X_i)_{\mathcal{F}_i, \mathcal{F}_i} HSIC(Y, Y)_{\mathcal{G}, \mathcal{G}}}}. \quad (6)$$

Based on Gretton et al. (2005a), an estimator of the HSIC can be computed from an  $N$  sample  $(x_i^j, y^j), j \in \{1, \dots, N\}$  of  $(X_i, Y)$ :

$$\widehat{HSIC}(X_i, Y)_{\mathcal{F}_i, \mathcal{G}} = \frac{1}{(N-1)^2} \text{Tr}(\mathbf{KHLH}), \quad (7)$$

where  $\mathbf{H} \in \mathbb{R}^{N \times N}$  is the centring matrix  $H_{ij} = \delta_{ij} - \frac{1}{N}$ , and  $\mathbf{K} \in \mathbb{R}^{N \times N}$  and  $\mathbf{L} \in \mathbb{R}^{N \times N}$  are the Gram matrices defined as  $K_{ij} = k_{X_i}(x_i, x_j)$  and  $L_{ij} = k_Y(y^i, y^j)$ , where  $k_{X_i}$  and  $k_Y$  are the kernel functions associated with each RKHS. In this

study, and following De Lozzo and Marrel (2014, 2016) and Da Veiga (2015), we choose a Gaussian kernel as it is a universal kernel that can fully characterize the independence of variables and that can be used for scalar or vectorial variables. For a vectorial variable  $\mathbf{x} \in \mathbb{R}^q$ , it is expressed as follows:

$$k(\mathbf{x}, \mathbf{x}') = \exp(-\lambda \|\mathbf{x} - \mathbf{x}'\|_2^2), \tag{8}$$

with  $\|\cdot\|_2$  the Euclidian norm in  $\mathbb{R}^q$  and where the hyperparameter  $\lambda$  is called the bandwidth parameter of the kernel. In this study, the bandwidth  $\lambda$  is estimated from the inverse of the empirical standard deviation of the sample.

When using a universal kernel, the HSIC indices can also be statistically used for screening purposes (De Lozzo and Marrel, 2014). A statistical test can be set with the null hypothesis “ $X_i$  and  $Y$  are independent”. Considering an experimental design of  $N$  points  $(x_i^1, \dots, x_i^N)$  and the associated output points  $(y^1, \dots, y^N)$ , an estimator  $\widehat{\text{HSIC}}(X_i, Y)$  of the dependence measure  $\text{HSIC}(X_i, Y)$  is firstly computed. Then  $B$  bootstrap versions  $\mathbf{Y}^{[1]}, \dots, \mathbf{Y}^{[B]}$  are resampled from the original output sample  $(y_1, \dots, y_N)$  with replacement so as to contain the same number of points  $N$ . For each  $\mathbf{Y}^{[b]}$  the input points associated with  $X_i$  are not resampled. Indeed, under the independence hypothesis, any values of  $Y$  can be associated with  $X_i$ . For each bootstrap version  $b$ , an estimator  $\widehat{\text{HSIC}}^{[b]}(X_i, Y)$  is computed. Then, the associated bootstrapped  $p$  value is given by

$$p\text{-val}_B = \frac{1}{B} \sum_{b=1}^B \mathbb{1}_{\widehat{\text{HSIC}}^{[b]}(X_i, Y) > \widehat{\text{HSIC}}(X_i, Y)}. \tag{9}$$

Finally, denoting  $\alpha$  the significance level, if  $p \text{ val}_B < \alpha$ , the independence hypothesis is rejected, otherwise it is accepted. In this study, such a statistical test is used to perform screening based on 100 bootstrap replicates and a 1 % significance level. The R code provided in De Lozzo and Marrel (2016) (see their Supplement) to compute the HSIC measure has been adapted in Python to perform both screening and ranking.

**2.4.3 Random forest**

Random forests (Breiman, 2001) belong to ensemble machine learning techniques. The method consists in creating a surrogate by averaging results from an ensemble of  $K$  decision trees created independently. A decision tree is composed of an ensemble of discriminatory binary conditions contained in nodes. Such conditions are hierarchically applied from a root node to a terminal node (tree leaf) (Rodriguez-Galiano et al., 2014). The input space is therefore successively partitioned into smaller groups that correspond to the nodes according to a response variable. Such splitting goes on until reaching a minimum threshold of members per node. In this study, we consider regression trees that focus on continuous response variables.

As each individual decision tree is very sensitive to the input dataset, bagging is used to avoid correlations between them and to ensure model stability. It consists in training each decision tree from a different training dataset smaller than the original one. Such subsets are built from the original one by resampling with replacement, making some members be used more than once, while others may not be used. Such a technique makes the random forests more robust when facing slight variations in the input space and increases accuracy of the prediction (Breiman, 1996, 2001). The samples that are not used to grow a tree are called “out-of-bag” (OOB) data and will be used for the test step. The RF workflow is summarized in Fig. 5.

RF structure can be used to provide knowledge about how influential each input factor is. This measure is referred to as *feature importance* in the RF formalism. The random forest is first trained on the targeted output variable  $Y$  using a  $N$ -point sample  $(\mathbf{X}^j, Y^j)$  for  $j \in \{1, \dots, N\}$ . Once the forest has been trained, each input factor  $X_i$  is permuted individually so as to break the link between  $X_i$  and  $Y$ . The effect of such permutation on the model accuracy is then investigated. A large decrease in accuracy indicates that the input factor is highly influential, whereas a small decrease in accuracy indicates that it has little influence. Different algorithms exist to compute such mean decrease in accuracy (MDA) (see Bénard et al., 2022, for an extensive review of the different formulations in the existing R and Python packages), and we focus here on the original formulation from Breiman paper (Breiman, 2001). The decrease in accuracy is originally computed from the mean square error between predictions from OOB data with and without permutation for each tree. Results are then averaged over all trees to get the MDA. The algorithm for feature importance calculation is extensively described in various papers (e.g. Soleimani, 2021; Bénard et al., 2022), and it is recalled in what follows.

1. For each tree  $k$ ,
  - estimate  $\hat{\epsilon}_{\text{OOB}_k}$ , the error from the OOB sample  $\mathcal{L}_k$ :

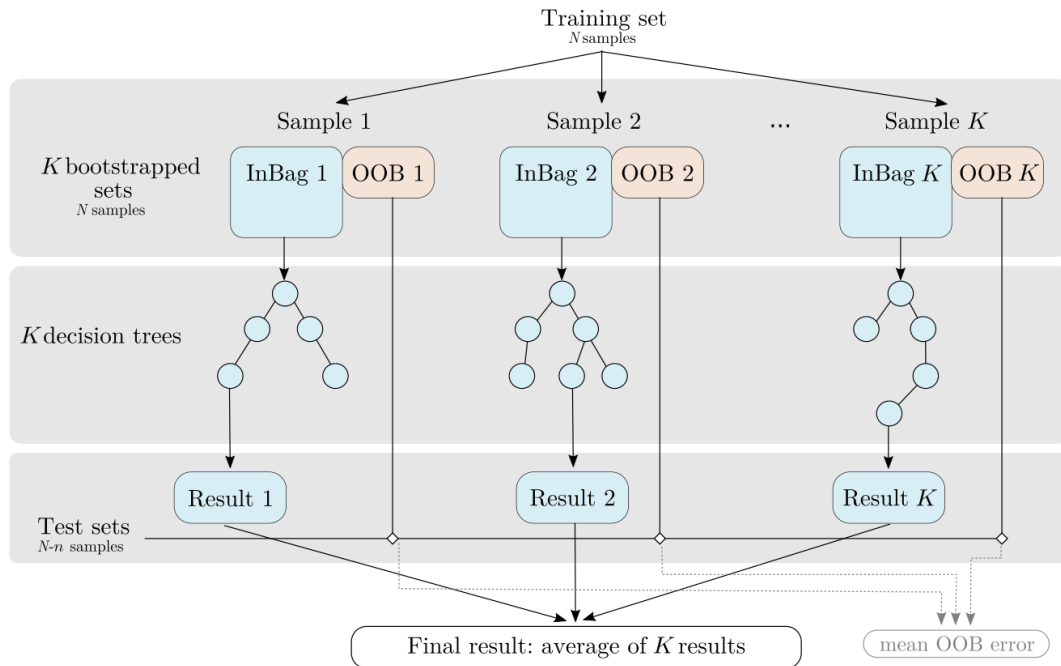
$$\hat{\epsilon}_{\text{OOB}_k} = \frac{1}{|\mathcal{L}_k|} \sum_{j \in \mathcal{L}_k} (y^j - \hat{\mathcal{M}}_{\text{RF}}(\mathbf{x}^j))^2, \tag{10}$$

where  $\hat{\mathcal{M}}_{\text{RF}}$  is the estimated RF metamodel.

- For each input factor  $X_i$ ,
  - randomly permute  $x_i$  in  $\{\mathbf{x}^j \in \mathcal{L}_k\}$  to generate a new input set  $\{\mathbf{x}^{j*} \in \mathcal{L}_k\}$ .
  - Estimate  $\hat{\epsilon}_{\text{OOB}_k}^*(i)$  using the permuted input set

$$\hat{\epsilon}_{\text{OOB}_k}^*(i) = \frac{1}{|\mathcal{L}_k|} \sum_{j \in \mathcal{L}_k} (y^j - \hat{\mathcal{M}}_{\text{RF}}(\mathbf{x}^{j*}))^2. \tag{11}$$

2. For each input factor  $X_i$ ,



**Figure 5.** RF workflow (adapted from Rodriguez-Galiano et al., 2014).  $K$  bootstrapped sets are extracted from the original training set. Part of each set “InBag” is used to grow an independent decision tree, and the final regression value is the average of all trees. The remaining portion of each tree’s “out-of-bag” (OOB) data is used as a test set.

- compute the mean decrease in accuracy  $MDA_i$  as

$$MDA_i = \frac{1}{K} \sum_{k=1}^K \hat{\epsilon}_{OOB_k} - \hat{\epsilon}_{OOB_k}^*(i), \quad (12)$$

where  $K$  is the total number of trees.

Despite the “black-box” aspect of RF building, recent works theoretically establish a link between mean decrease in accuracy and Sobol’ total indices when input parameters are assumed independent. Indeed, Gregorutti et al. (2017) establish that for all input parameters  $X_j$ ,

$$ST_i = \frac{MDA_i}{2\text{Var}[Y]}. \quad (13)$$

In this study, the randomForestSRC R package (Ishwaran and Kogalur, 2020) is used to obtain feature importance measures, and the number of trees used to train the RF is set to 500.

## 2.5 Robustness and reliability assessment of the sensitivity indices

In order to assess the robustness of the calculated sensitivity indices, an additional 200-point test set is first used to assess PCE and RF metamodel performances. In a second time, 95 % confidence intervals on sensitivity indices are computed for all methods but by different means: the bootstrap resampling procedure provided in UQLab is used to calculate confidence intervals on PCE-based Sobol’ indices (see Marelli

and Sudret, 2018, for justification). In the case of random forests, building extra bootstraps could affect the consistency of the OOB sample as the RF structure already includes a bootstrap step to build each tree. Then, we use a subsampling approach without replacement with a subsample size set to 80 % of the initial sample size to get error bounds on feature importance measures. The same procedure is applied to estimate error bounds on HSIC indices. As typically found in similar studies (Archer et al., 1997; Yang, 2011), 1000 bootstrap resamples are used for all methods. In addition, the reliability of each ranking method is assessed by monitoring its convergence properties. To do so, sensitivity indices and associated confidence intervals are calculated on samples of growing sizes. A new sample is generated for each size in [50, 100, 250, 750, 1000, 2000], where the maximum size (2000) points corresponds to the maximum computational budget available for ranking we guess for a catchment-scale application. It is noted that we create a new sample for each sample size to keep all samples as independent as possible, but such an approach may be too costly in a catchment scale application. In such a case, another strategy could be to generate the sample for ranking from the initial LHS used for screening, dropping the dimensions corresponding to the non-influential inputs.

## 2.6 Aggregated sensitivity indices

After computing local sensitivity indices for each landscape unit on a scalar quantity, aggregated indices are computed at

the catchment scale (step 2b, Fig. 4). Catchment-scale sensitivity indices are computed considering a multidimensional output  $Y \in \mathbb{R}^d$  that gathers scalar outputs for each landscape unit. Sobol' indices are aggregated at the catchment scale following the formulation by Gamboa et al. (2013) for generalized Sobol' indices:

$$S_u = \frac{\sum_{j=1}^d \text{Var}[Y_j] S_{u,j}}{\sum_{j=1}^d \text{Var}[Y_j]}, \quad (14)$$

where  $u$  is a subset of  $\{1, \dots, M\}$ ,  $\text{Var}[Y_j]$  is the variance of the scalar  $j$ th component of  $Y$  and  $S_{u,j}$  is the Sobol' indices of subset  $u$  on  $Y_j$ . Equation (14) then formulates catchment-scale indices as an average of Sobol' indices on each landscape unit weighted by local output variances. First- and total-order Sobol' indices can be notably computed this way. Their computation can be made in a second step after performing local Sobol' analysis on each landscape unit, but a direct estimator is also proposed in Gamboa et al. (2013) avoiding numerous local analysis in the case of high-dimensional model output. For HSIC and RF sensitivity indices, the definitions for scalar output remain valid for vectorial output. Catchment-scale indices can thus be directly computed on  $Y$  when moving to a multidimensional case.

### 3 Results

#### 3.1 Screening

Screening is performed on the 4000-point LHS. Simulations are run on the HIICS cluster (26 nodes, 692 cores, 64 to 256 GO of RAM per server) available at INRAE, France, for a simulation time of 12 000 CPU hours. Screening is performed at a site scale, on each HU individually, to remain as conservative as possible. Influential parameters at the catchment scale are then deduced from the union of influential parameters for each site. After screening and union, 42 influential parameters are selected for water subsurface flow, 54 parameters are selected for pesticide subsurface flow, 43 parameters are selected for water surface runoff, and 45 parameters are retained for pesticide surface runoff. The remaining parameters are given in Appendix C1.

#### 3.2 Ranking on a single HU

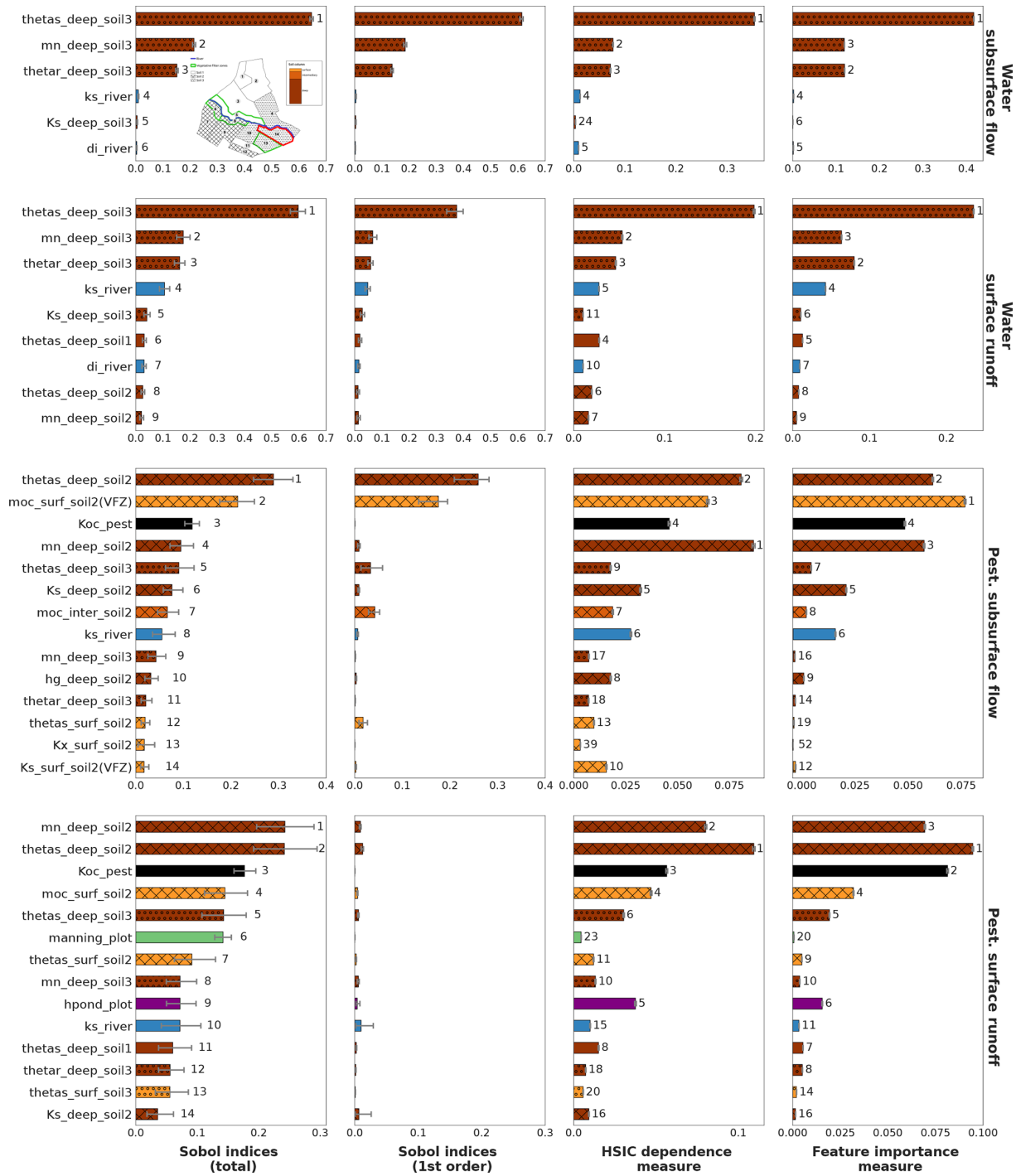
For the ranking task, the three methods (Sobol' indices from PCE, HSIC and RF) are applied on each HU based on a 2000-point LHS generated for each scrutinized variable from the set of screened input parameters. Again, simulations are performed on the HIICS cluster, for a simulation time of 6000 CPU hours per variable. Columns 1 and 2 in Fig. 6 show Sobol' total- and first-order indices, while columns 3 and 4 show HSIC and RF sensitivity indices for the most influential parameters according to Sobol' total-order indices for the four variables on HU14.

#### 3.2.1 Physical interpretation

Considering Sobol' indices, influential parameters that are identified highly differ from one output variable to the other. They are linked to distinct physical processes that may interact with the other ones. This way, sensitivity analysis brings knowledge about the way PESHMELBA represents the hydrological functioning of the virtual catchment. Water subsurface flow (top line) is driven by deep soil hydrodynamic parameters both related to vertical infiltration and subsurface saturated transfers. Water surface runoff (line 2) is also mainly influenced by deep soil parameters. Overland flow is therefore identified as being mostly due to saturation rather than to rainfall excess. Subsurface exchanges with the river are also identified as an influential process as the river bed saturated conductivity ( $Ks_{river}$ ) is part of the most influential parameters. Such a finding is consistent with the position of HU14, which is directly connected to the river but also to many plots (see Fig. 1). Pesticide variables (line 3 and 4) are influenced by a higher diversity of parameters that characterize contrasted and interactive physical processes. Indeed, both pesticide subsurface flow and surface runoff are mostly influenced by deep soil parameters such as the saturated water content ( $\theta$ ). Parameters linked to pesticide adsorption such as the pesticide adsorption coefficient and the organic carbon content ( $OC$ ) of surface and intermediary soil horizons also rank among the top influential parameters. Additionally, the roughness coefficient ( $n_{plot}$ ) and the ponding height ( $h_{pond\_plot}$ ) that are related to surface runoff calculation are also ranked as highly influential on pesticide surface runoff.

#### 3.2.2 Comparison of methods

The rankings that are provided by the three methods are broadly consistent giving confidence in their robustness. Looking in more detail, HSIC and RF indices are more similar to Sobol' total-order indices than to Sobol' first-order indices, contrary to the conclusions of De Lozzo and Marrel (2016) on comparison of Sobol' and HSIC indices. The most influential parameters that are identified by Sobol' total-order indices are also captured by the other methods. Indeed, the different rankings show that the top-10 rankings based on Sobol' total-order indices contain at least the five most influential parameters based on HSIC and RF rankings. Then, Fig. 6 does not miss any preponderant parameters for the HSIC and RFs. In addition, while rankings from the three methods only exhibit slight differences for water variables, differences between rankings are more pronounced for pesticide variables. In that case, most parameters have zero or very low first-order Sobol' indices, characterizing nearly purely interacting effects. However, not all differences between rankings come from interactive effects. As an example,  $n_{plot}$  and  $h_{pond\_plot}$  that appear in the pesticide surface runoff top-10 ranking according to Sobol' in-



**Figure 6.** Sobol’ total- and first-order indices computed using PCE, HSIC and RF site sensitivity indices for all output variables on HU14 with associated 95 % confidence intervals. RF feature importance measures are normalized by  $2\text{Var}[Y]$  following Eq. (13). HU14 is displayed with a red contour in the top left figure. For all methods, displayed parameters are the most influential parameters regarding Sobol’ total-order indices. The bar colours are related to physical processes: brown is related to soil parameters, and the darker the brown, the deeper the parameter; blue is related to river parameters, whereas green is related to vegetation parameters. The filling in of brown bars refers to the soil type of the parameter: soil 1 is not filled, and soil 2 is cross-hatched, whereas soil 3 is filled with circles. For each method, numbers next to bars stand for the ranking position of parameters.

**Table 4.**  $Q_2$  score for all variables on HU14 calculated from the 200-point test set. The  $Q_2$  score is calculated as  $Q^2 = 1 - \frac{\sum_{i=1}^N (\mathcal{M}(X^i) - Y^i)^2}{\sum_{i=1}^N (Y^i - \bar{Y})^2}$ , where  $\bar{Y} = \frac{1}{N} \sum_{i=1}^N Y^i$  is the empirical mean of the sample.

	PCE	RF
WaterLateralFlow	0.98	0.85
WaterSurfaceRunoff	0.80	0.44
PesticideLateralFlow	0.75	0.55
PesticideSurfaceRunoff	0.75	0.51

indices are both characterized by nearly fully interactive effects. However, `hpond_plot` is also ranked as influential according to the HSIC and RFs, while `manning_plot` is missed by both of them.

Regarding error bounds, they are very small for the HSIC and RFs, contrary to Sobol' indices estimates. The HSIC is expected to be very accurate from very small sample sizes (Da Veiga et al., 2021), which may explain such differences of magnitude for error of Sobol' and HSIC indices. Regarding RF error bounds, we incriminate the resampling strategies that differ from the Sobol' method. Indeed, while the bootstrap technique has been proven to assess the quality of the PCE (Marelli and Sudret, 2018), the subsampling technique set on RF only targets the precision of feature importance measures. A more adapted subsampling approach, for example, based on Ishwaran and Lu (2019) should probably be further investigated on bigger samples to better compute RF sensitivity indices and to accurately assess their quality. In addition to Fig. 6, Table 4 gathers the quality scores  $Q_2$  for PCE and RF metamodels on the test set. Results show that the RF metamodel performs much more poorly than the PCE for all variables. Such poor performances may explain the discrepancies between RF feature importance measures and Sobol' total-order indices. Indeed, RF feature importance indices have been proven to relate to Sobol' total-order indices (Gregorutti et al., 2017), but this relation (see Eq. 13) is not respected in this study. These results thus underline how crucial it is to analyse both the quality of the metamodel and the quality of the indices it calculates and not just one of the two.

### 3.2.3 Convergence rate

In order to assess each method convergence rate, sensitivity indices are calculated for growing sample sizes, from 50 to 2000 points. Results for water surface runoff and pesticide surface runoff are presented respectively in Figs. 7 and 8. Results for water (i.e. pesticide) subsurface flow are not presented as they come to the same conclusions as for water (i.e. pesticide) surface runoff.

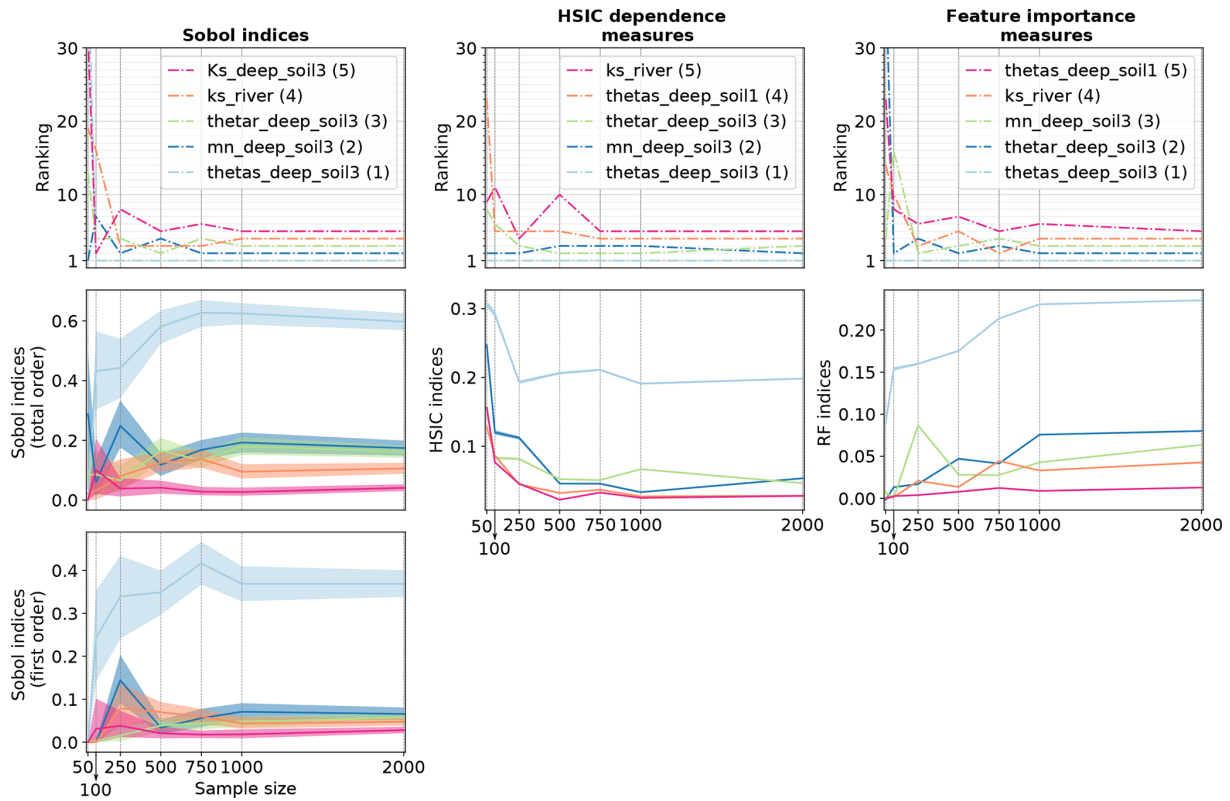
Figure 7 shows that Sobol' indices are close from stability for sample sizes from 1000 points as ranking and Sobol' total- and first-order indices are rather stable. The 95 % con-

fidence intervals associated with Sobol' indices (first- and total-order) also seem to be rather stable for sample sizes from 1000 points. Considering HSIC and RF indices, ranking differences for the five most sensitive input factors are less than one position for sample sizes from 1000 points, which roughly suits the convergence criteria for parameter ranking defined in Sarrazin et al. (2016). Identically, for both methods, sensitivity indices' values and 95 % confidence intervals show little variability. As already noticed in Fig. 6, the 95 % confidence intervals for the HSIC and RFs are much smaller than for PCE. Again, such results should be interpreted with caution as confidence intervals are not calculated in the same way as for the different methods. On the whole, the HSIC method reaches convergence first as ranking and sensitivity indices' values stabilize for sample sizes from 750, while 1000 points are necessary for Sobol' and RF methods.

Results about pesticide surface runoff (Fig. 8) are more contrasted. Ranking and sensitivity indices' values from PCE show more variability than for water variables. This is especially the case for first-order Sobol' indices, which shows that 2000 may not be a sufficient sample size to get robust results. Ranking and total-order indices are more stable, except for the parameter `Koc_pest`. This parameter is characterized by fully interactive effects, which may explain why calculating robust Sobol' indices is a more complex task. Ranking and sensitivity measures from the HSIC stabilize from 750 points, while the error bounds remain very limited for all sample sizes. Finally, RF ranking and indices are almost stable from 1000 points, except for `Koc_pest`, whose RF value still shows variability for the biggest sample sizes.

### 3.3 Landscape analysis

In this section, we focus on Sobol' indices (first- and total-order) despite larger error bounds as it is the only method used in this study that allows us to get separate information on interactive effects. Site rankings such as those presented in Fig. 6 are gathered for all HUs in the form of sensitivity maps in Fig. 9 for water surface runoff and in Fig. 10 for pesticide surface runoff. Broadly speaking, both maps show strong spatial heterogeneities regarding influential parameters, and a contrasted behaviour between right and left banks can be identified. For both output variables, hydrodynamic parameters (thetas, `thetar` and `mn`) of deep horizon from soil 1 (i.e. 2 and 3) are mainly influential only on HUs characterized by soil 1 (i.e. 2 and 3) (see Fig. 2 for a reminder of soil types). The local hydrodynamic parameters are found to be dominating, explaining the output variable variance. A particular case is HU4 (indicated by an array on both Figs. 9 and 10), which is characterized by soil 3, while parameters from soil 1 explained most of the variance of both water and pesticide surface runoff variables. The location of HU4, near the outlet, downstream several soil-1 HUs, may explain such spatial interactions. In addition to specific soil parameters, other parameters such as Manning's roughness on vineyard plots



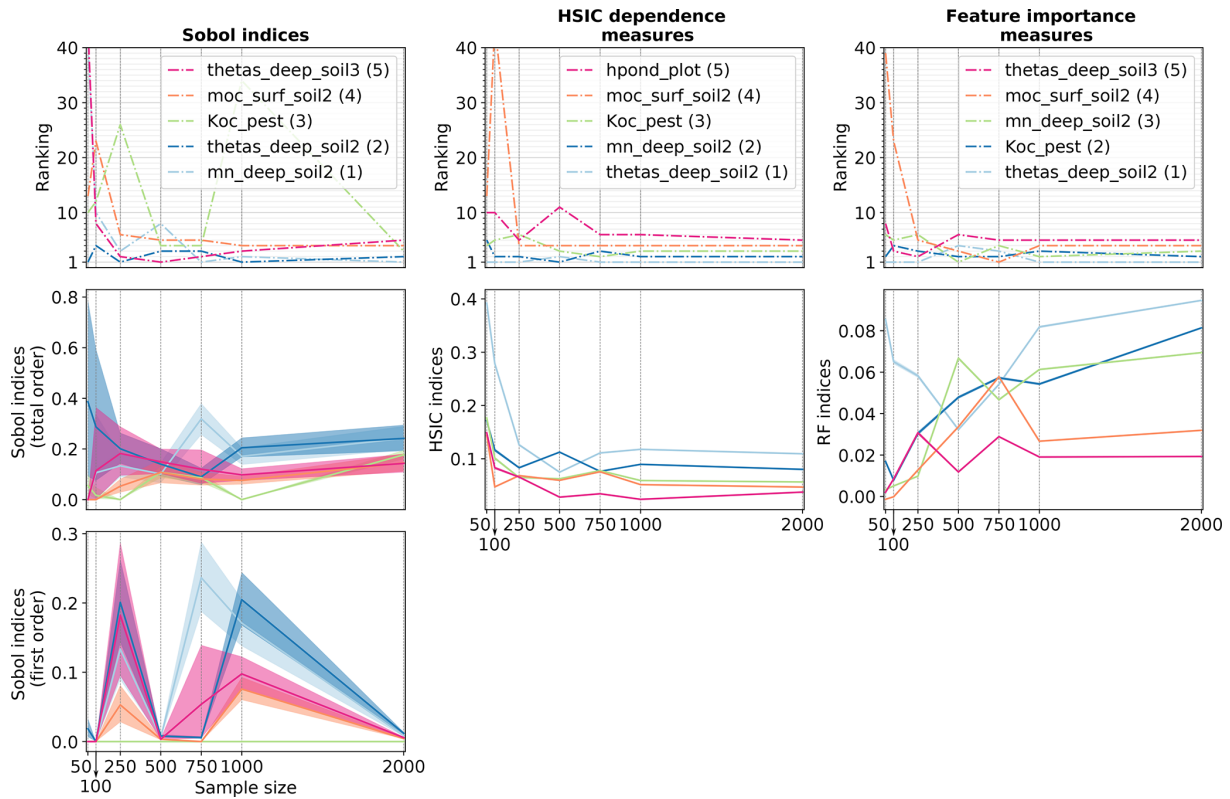
**Figure 7.** Convergence plot for solute water runoff variable: the figure shows the ranking for the five most influential parameters identified with a sample of 2000 points as well as the corresponding sensitivity indices’ values with associated 95 % confidence intervals for growing sample sizes (even if they are hardly visible for the HSIC and RFs, confidence intervals are displayed on all plots from line 2). In the legend, the number in brackets is the ranking position of the parameter in the case of a sample of 2000 points.

(manning\_plot) or the coefficient of adsorption (Koc\_pest) have a greater influence on HUs from the right bank (bottom part of the catchment in the figure). In addition, comparison of first-order and total-order maps shows quite similar results for water surface runoff on the one hand. It indicates that direct effects are significant for all influential parameters. On the other hand, direct effects are far from dominant on pesticide surface runoff. Once again, most parameters are influential nearly only in interaction with other parameters since the first-order indices are very low compared to the total-order indices.

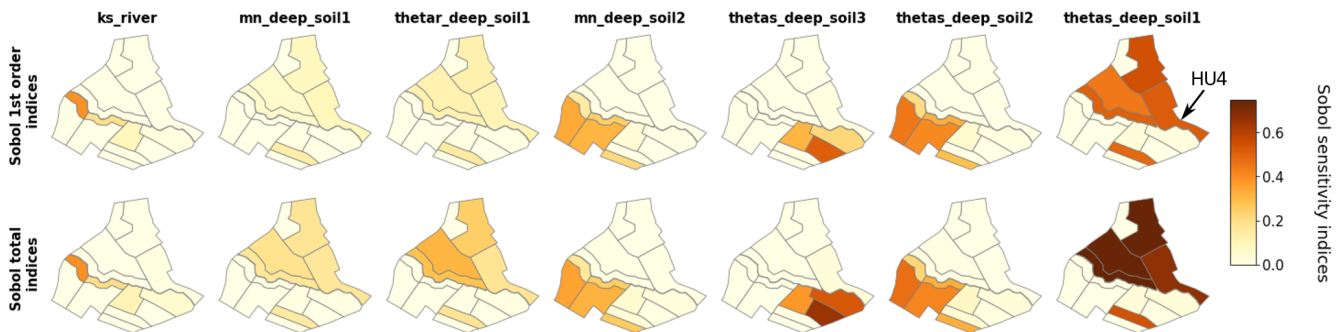
Finally, Fig. 11 shows aggregated sensitivity indices for water and pesticide surface runoff variables following Gamboa et al. (2013). Since the two banks have contrasted behaviours, aggregated indices are first calculated at the intermediary scale of the bank, then at the catchment scale. For both output variables, rankings strongly differ on each slope. As proposed at a local scale in Sect. 3.2, aggregated indices at this scale may constitute summarized information about the physical processes dominating in PESHMELBA to explain the output variable. For water surface runoff, hydrodynamic soil parameters related to vertical infiltration (thetas, thetar and mn) dominate. The influence of Ks\_river is only signif-

icant in the right bank. The difference of altitude between the right and left bank may explain this contrast in the activation of saturated exchanges between water tables and the river. For pesticide surface runoff, deep horizon parameters from soil 1 and the pesticide adsorption coefficient (Koc) explain a major portion of the output variance on the left bank, while pesticide half-life time (DT50) and surface runoff parameter (hpond\_plot) have lower or no impact. Conversely, surface parameters (manning\_plot and hpond\_plot) have a higher impact on the right slope. In that bank, soil horizons are characterized by lower permeabilities that may result in stronger surface runoff generation than on left bank. In addition, pesticide parameters (Koc and DT50) are also more influential. More broadly, these results show that pesticide surface runoff may result from the activation and interactions of more physical processes on the right bank than on the left bank.





**Figure 8.** Convergence plot for solute surface runoff variable: the figure shows the ranking for the five most influential parameters identified with a sample of 2000 points as well as the corresponding sensitivity indices' values with associated 95 % confidence intervals for growing sample sizes (even if they are hardly visible for the HSIC and RFs, confidence intervals are displayed on all plots from line 2). In the legend, the number in brackets is the ranking position of the parameter in the case of a sample of 2000 points.



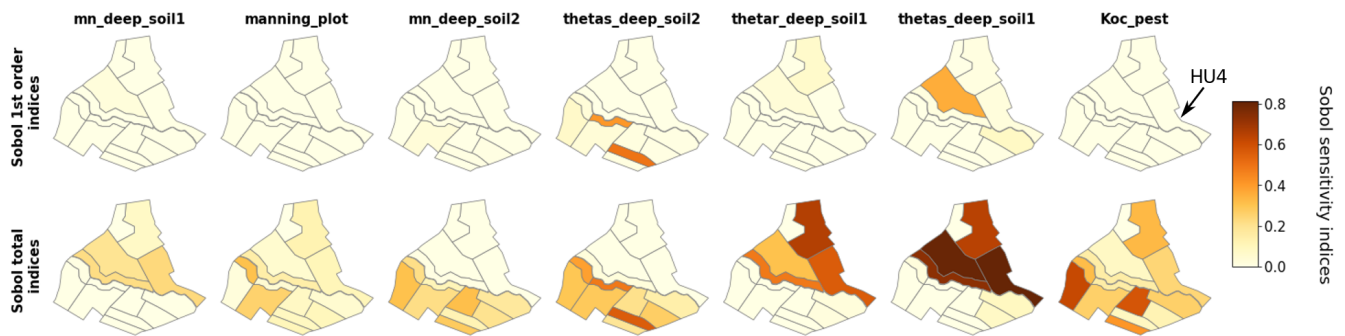
**Figure 9.** Maps of Sobol' site sensitivity indices for water surface runoff for the most significantly influential parameters.

## 4 Discussion

### 4.1 On screening

For all variables considered, the number of input parameters retained after screening remained quite high, proving that performing screening on PESHMELBA variables is a challenging task. We can first incriminate the many physical processes interacting in PESHMELBA in a spatially distributed way, each of them with its own set of characteristic param-

eters. However, it can also be explained by the methodology that may not be discriminating enough. Many previous studies developed efficient screening techniques for complex environmental models (e.g. Tang et al., 2007; Nossent et al., 2011; Touzani and Busby, 2014; Becker et al., 2018; Garcia et al., 2019; Sheikholeslami et al., 2019). Exploring other screening approaches is beyond the scope of this study, but future work may focus on applying and critically comparing these techniques, especially with very limited sample size (inferior to 2000 points for example).



**Figure 10.** Maps of Sobol' site sensitivity indices for pesticide surface runoff for the most significantly influential parameters.

#### 4.2 On the choice of a ranking method

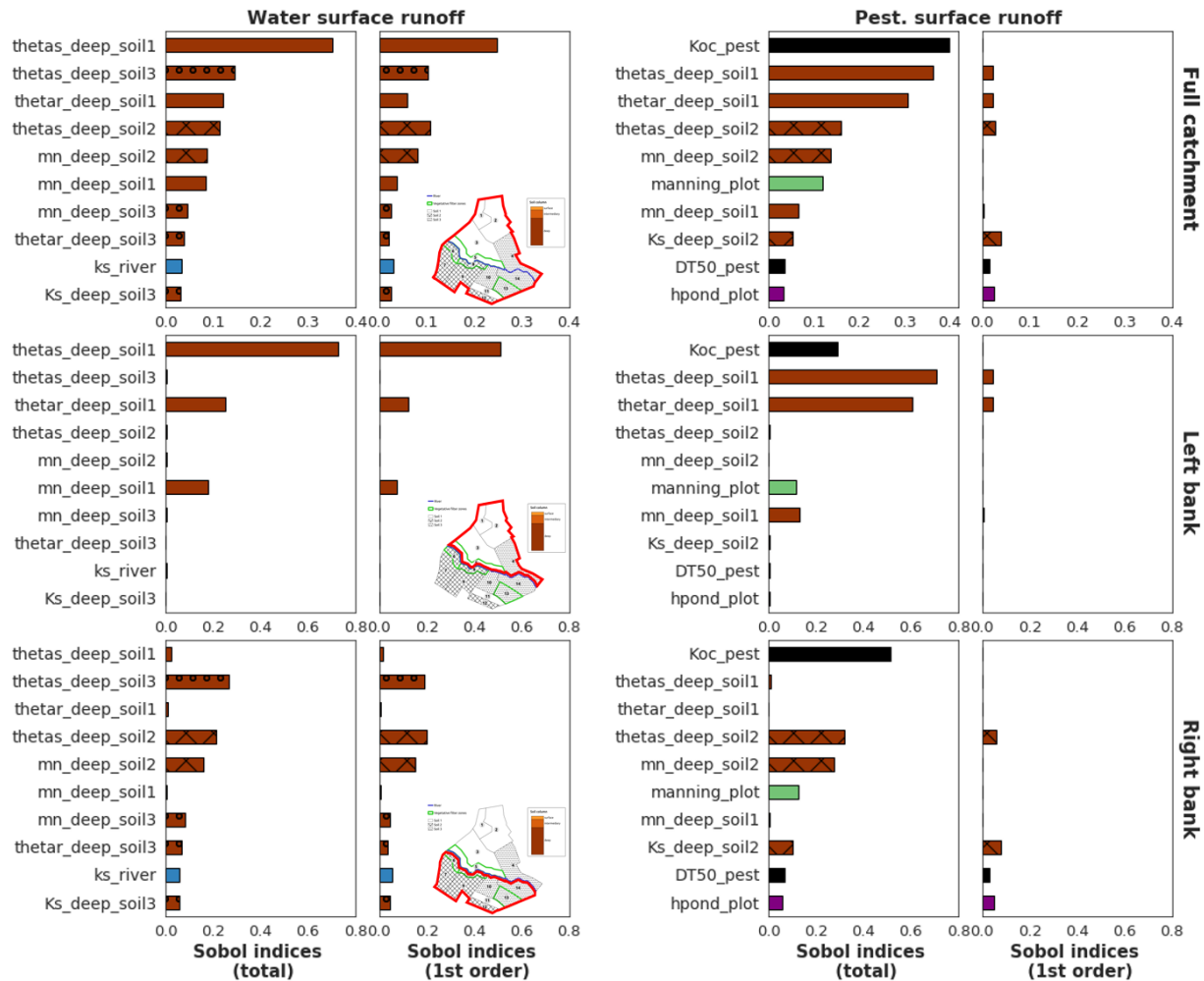
By exploring several methods for ranking, the aim was to analyse their specificities and the interest of each one for the sensitivity analysis of a complex environmental model, characterized by many parameters and a high computational cost such as PESHMELBA. Considering the results of this study, we believe that the choice of the method depends on the properties of the model, the objective of the sensitivity analysis and the sample size available:

1. The Sobol' method remains attractive when sensitivity analysis is used to gain knowledge about the model by finely analysing its behaviour. Indeed, Sobol' indices provide a clear interpretation of the calculated indices (percentages of variance explained) and explicit information about the interactions between parameters. These elements are particularly valuable when one wishes to use sensitivity analysis to understand the functioning of the model, and this is why this approach is still widely used in the hydrological community. In the case of variables that are reasonably complex and that are not characterized by too much interactions of physical processes (such as water variables in our case), using chaos polynomials to estimate Sobol' indices is particularly interesting and efficient since it allows the use of a pre-existing sample, of very limited size. Conclusions are much contrasted for complex variables such as pesticide variables as convergence results showed that 2000 points may not be enough to get fully robust Sobol' indices. This is particularly the case for parameters that are mainly characterized by interactive effects.
2. If the sensitivity analysis aims at simplifying the model or focusing the calibration efforts, if the physical interpretation of the results is not a priority and if one has a pre-existing sample of very limited size (inferior to 750 points in our case), the use of HSIC indices is a good option as it provides robust sensitivity indices. However, it is important to note that using the HSIC dependence measure for sensitivity analysis is a recent idea and that there is still little knowledge available about identify-

ing and differentiating the types of dependency that are captured. In addition, the choice of the kernel may affect the ranking results because each specific kernel is likely to give more or less importance to the infinite number of dependency forms that are captured by the HSIC. The question of the choice of the kernel is delicate, and it is still not addressed very much in the literature. While a few papers propose to choose the type and parameterization of the kernels in a way that maximizes the possible dependence between  $X_i$  and  $Y$  (Fukumizu et al., 2008; Balasubramanian et al., 2013), the interpretation of the results seems to be less clear. On the other hand, there are still relatively few works that apply this method for GSA, and the limitations are not necessarily all identified yet. Using the HSIC for a classification exercise will therefore remain delicate as long as there is no consensus on the choice of the kernel and the interpretation of the results. However, these problems do not arise when using the HSIC for screening, and such a method is therefore perhaps to be preferred for this type of exercise.

3. The RF indices are also of interest in sensitivity analysis task as they are supposed to provide an estimator of total-order Sobol' indices. Those indices can thus be easily interpreted, and as for the other methods, they can be estimated from a pre-existing sample. However, PCE is still to be preferred since it provides more complete information including not only the total-order Sobol' indices but also the indices at all orders. In addition, our results showed that the metamodel constructed by RF is of lower quality than the one constructed by PCE at equivalent sample size, giving less confidence in the resulting sensitivity indices.

Beyond the comparison of the different methods, we also tried to evaluate if it was possible and useful to combine several of these methods. However, considering the results obtained, we believe that combining the tested methods is still of little interest for hydrologists to better understand the model functioning. Indeed, the differences we found in rankings remain difficult to interpret. This is particularly the case



**Figure 11.** Sobol' first-order and total-order aggregated sensitivity indices for water surface runoff (left) and pesticide surface runoff (right) calculated at the scale of the catchment (top), left bank (middle) and right bank (bottom). Displayed parameters are the 11 most influential parameters regarding Sobol' indices at the catchment scale for each output variable. The bar colours are related to physical processes: brown is related to soil parameters, and the darker the brown, the deeper the parameter is; blue is related to river parameters, and green is related to vegetation parameters. The filling in of brown bars refers to the soil type of the parameter: soil 1 is not filled, and soil 2 is cross-hatched, while soil 3 is filled with circles.

when combining Sobol' and HSIC indices, due to the fact that the results from the HSIC dependence measure remain fuzzy to interpret.

### 4.3 On choosing local or aggregated sensitivity analysis

Results about landscape analysis showed that, on the one hand, sensitivity maps provide local, detailed information about influential parameters on each location of the catchment. However they are computationally costly as one GSA per HU must be performed. This approach may be hard or even impossible to transpose to a real catchment scale composed of several hundred elements. On the other hand, catchment-scale aggregated indices provide synthetic information at a lower computational cost, but the spatialized aspect of the GSA is lost. As pointed out in Marrel et al. (2015),

both approaches are complementary and provide precious knowledge about the model functioning, but they cannot always be performed together. As a compromise, we propose to use the scale of the bank, or more generally of the hill-slope, as it may constitute an adapted intermediary scale to meet both requirements of detailed results for physical interpretation and computational efficiency. Indeed, in this study Sobol' aggregated indices were directly computed from site sensitivity indices as they were available, but a pick-freeze estimator (Gamboa et al., 2013; De Lozzo and Marrel, 2016) can be used for a direct computation of Sobol' generalized indices. In our case, such overview of sensitivity analysis allows us to focus calibration efforts on deep soil hydrodynamic parameters and pesticide adsorption coefficient to im-

prove the quality of the simulation of both water and pesticide surface flows.

## 5 Conclusions

In this paper, we have described the first global sensitivity analysis of the modular and coupled PESHMELBA model. For this experiment, a simplified catchment was set in order to explore different approaches for GSA and to propose a methodology for future real applications. First, we performed screening using an independence test based on the HSIC dependence measure, dividing the dimension of the problem by 3. Second, we compared several innovative methods to compute sensitivity measures on each landscape element individually. Sobol' indices were found to be particularly attractive as they provide easy-to-interpret sensitivity measures. However, in the case of complex variables with dominant interactive effects, results showed that they may not be computed from very small samples. Third, we gathered such local sensitivity indices into sensitivity maps that highlighted local contributions of parameters. Finally, we computed aggregated indices at larger scales, on the whole catchment and on each bank hillslope since this scale still reflects spatial heterogeneities of hydrodynamic processes. This study constitutes the first attempt of global sensitivity analysis of the PESHMELBA model. Future research should go a step further by considering the other sources of uncertainties that can affect the model and interact with parameter uncertainties. The impact of forcings, soil types, quantities and dates of application of pesticides should be addressed as already done in Holvoet et al. (2005) for instance. Additionally, parameters were assumed to be independent in this study, but this assumption may be arguable, especially for hydrodynamic parameters. For the three methods, sensitivity indices that are produced are meaningless in the case of dependent inputs. Dealing with dependent parameters has already been explored in the case of Sobol' indices and the HSIC measure, for example, based on Shapley effects (Da Veiga et al., 2021), but again, these formulations should be extensively tested with complex variables and very small sample sizes. Finally, it would also be necessary to investigate sensitivity of some time series to get a more comprehensive vision of the model functioning. To do so, the temporal series can be analysed as a multivariate output for example with clustering-based GSA (Roux et al., 2021) or using the principal components of the model's functional outputs. The definition and the use of adequate hydrological signatures such as proposed in Branger and McMillan (2020) and Horner (2020) may also be of interest to understand space–time variability and to capture a broader range of physical processes. Global sensitivity analysis is a necessary but not yet systematic step to model evaluation, especially in the case of spatialized, risk assessment models that can be complex to deal with. This study brings additional knowledge on GSA strategies for modellers who

deal with such complex models and thus paves the way for systematic analysis of environmental exposure models.

## Appendix A: Parameters for LAI evolution law

**Table A1.** Top: parameters and associated dates used to describe LAI evolution for vineyard cover. Bottom: constant LAI value set on grassland cover.

Parameter	Value	Date
Vineyard (plot)		
LAI <sub>min</sub> [–]	0.01	1 February
LAI <sub>max</sub> [–]	2.5	1 May
LAI <sub>harv</sub> [–]	0.01	10 November
Grassland (VFS)		
LAI [–]	5	–

Appendix B: Input parameter distributions

**Table B1.** Distribution and statistics of the assigned probability density functions (pdfs) for the 145 input parameters, uniform: U(min,max), triangular: T(min,mean,max), normal: N(mean,standard deviation), lognormal: LN(mean,standard deviation) and truncated normal: TN(mean, standard deviation, min,max).

Soil parameters		
soilhorizon_thetas_2	[m <sup>3</sup> m <sup>-3</sup> ]	N(0.3362, 0.00336)
soilhorizon_thetar_2	[m <sup>3</sup> m <sup>-3</sup> ]	TN(0.0510, 0.0128, 0, 1)
soilhorizon_Ks_2	[m s <sup>-1</sup> ]	LN(9.5e-06, 1.97e-05)
soilhorizon_hg_2	[m]	N(-0.0329, 0.00329)
soilhorizon_mn_2	[-]	N(0.1988, 0.0199)
soilhorizon_Ko_2	[m s <sup>-1</sup> ]	LN(-7e-06,5.5e-07)
soilhorizon_L_2	[-]	U(-7.8216, -5.2144)
soilhorizon_bd_2	[g cm <sup>-3</sup> ]	U(1.1768, 1.7652)
soilhorizon_moc_2	[g g <sup>-1</sup> ]	U(0.0024, 0.0054)
soilhorizon_thetas_3	[m <sup>3</sup> m <sup>-3</sup> ]	N(0.3202, 0.0320)
soilhorizon_thetar_3	[m <sup>3</sup> m <sup>-3</sup> ]	TN(0.0812, 0.0203, 0, 1)
soilhorizon_Ks_3	[m s <sup>-1</sup> ]	LN(8.18e-06, 5.5e-07)
soilhorizon_hg_3	[m]	N(-0.0209, 0.00209)
soilhorizon_mn_3	[-]	N(0.2046, 0.0205)
soilhorizon_Ko_3	[m s <sup>-1</sup> ]	LN(-3.7e-06, 5.5e-07)
soilhorizon_L_3	[-]	U(-5.0844, -3.3896)
soilhorizon_bd_3	[g cm <sup>-3</sup> ]	U(1.2536, 1.8804)
soilhorizon_moc_3	[g g <sup>-1</sup> ]	U(0.0006, 0.0014)
soilhorizon_thetas_4	[m <sup>3</sup> m <sup>-3</sup> ]	N(0.2844, 0.0284)
soilhorizon_thetar_4	[m <sup>3</sup> m <sup>-3</sup> ]	TN(0.0661, 0.0165, 0, 1)
soilhorizon_Ks_4	[m s <sup>-1</sup> ]	LN(6.65e-06, 5.5e-07)
soilhorizon_hg_4	[m]	N(-0.0599, 0.00599)
soilhorizon_mn_4	[-]	N(0.2274, 0.0227)
soilhorizon_Ko_4	[m s <sup>-1</sup> ]	LN(-1.82e-06, 5.5e-07)
soilhorizon_L_4	[-]	U(-0.1716, -0.1144)
soilhorizon_bd_4	[g cm <sup>-3</sup> ]	U(1.2240, 1.8360)
soilhorizon_moc_4	[g g <sup>-1</sup> ]	U(4.3840 10 <sup>-4</sup> , 9.6160 10 <sup>-3</sup> )
soilhorizon_thetas_6	[m <sup>3</sup> m <sup>-3</sup> ]	N(0.3537, 0.0354)
soilhorizon_thetar_6	[m <sup>3</sup> m <sup>-3</sup> ]	TN(0, 0.0093, 0, 1)
soilhorizon_Ks_6	[m s <sup>-1</sup> ]	LN(5.64e-06, 5.5e-07)
soilhorizon_hg_6	[m]	N(-0.066, 0.0066)
soilhorizon_mn_6	[-]	N(0.1289, 0.0129)
soilhorizon_Ko_6	[m s <sup>-1</sup> ]	LN(-6.06e-06, 5.5e-07)
soilhorizon_L_6	[-]	U(7.7240, 19.3100)
soilhorizon_bd_6	[g cm <sup>-3</sup> ]	U(1.2704, 1.9056)
soilhorizon_moc_6	[g g <sup>-1</sup> ]	U(0.0042, 0.0094)
soilhorizon_thetas_7	[m <sup>3</sup> m <sup>-3</sup> ]	N(0.3247, 0.0325)
soilhorizon_thetar_7	[m <sup>3</sup> m <sup>-3</sup> ]	TN(0, 0.0093, 0, 1)
soilhorizon_Ks_7	[m s <sup>-1</sup> ]	LN(3.39e-06, 5.5e-07)
soilhorizon_hg_7	[m]	N(-0.0718, 0.00718)
soilhorizon_mn_7	[-]	N(0.0751, 0.0075)
soilhorizon_Ko_7	[m s <sup>-1</sup> ]	LN(-7.87e-06, 5.5e-07)
soilhorizon_L_7	[-]	U(-12, -8)
soilhorizon_bd_7	[g cm <sup>-3</sup> ]	U(1.3256, 1.9884)
soilhorizon_moc_7	[g g <sup>-1</sup> ]	U(0.0019, 0.0051)
soilhorizon_thetas_8	[m <sup>3</sup> m <sup>-3</sup> ]	N(0.4162, 0.0416)
soilhorizon_thetar_8	[m <sup>3</sup> m <sup>-3</sup> ]	TN(0, 0.0093, 0, 1)
soilhorizon_Ks_8	[m s <sup>-1</sup> ]	LN(9.42e-07, 5.5e-07)
soilhorizon_hg_8	[m]	N(-0.3018, 0.03018)
soilhorizon_mn_8	[-]	N(0.10000, 0.0100)
soilhorizon_Ko_8	[m s <sup>-1</sup> ]	LN(-9.37e-06, 5.5e-07)
soilhorizon_L_8	[-]	U(8, 20)
soilhorizon_bd_8	[g cm <sup>-3</sup> ]	U(1.2304, 1.8456)
soilhorizon_moc_8	[g g <sup>-1</sup> ]	U(0.0018, 0.0037)

Table B1. Continued.

Soil parameters		
soilhorizon_thetas_9	[m <sup>3</sup> m <sup>-3</sup> ]	N(0.3322, 0.0332)
soilhorizon_thetar_9	[m <sup>3</sup> m <sup>-3</sup> ]	TN(0.0770, 0.0192, 0, 1)
soilhorizon_Ks_9	[m s <sup>-1</sup> ]	LN(6.6e-06, 5.5e-07)
soilhorizon_hg_9	[m]	N(-0.0671, 0.00671)
soilhorizon_mn_9	[-]	N(0.2582, 0.0258)
soilhorizon_Ko_9	[m s <sup>-1</sup> ]	LN(-5.92e-06, 5.5e-07)
soilhorizon_L_9	[-]	U(0.3376, 0.8440)
soilhorizon_bd_9	[g cm <sup>-3</sup> ]	U(1.1664, 1.7496)
soilhorizon_moc_9	[g g <sup>-1</sup> ]	U(0.0023, 0.0051)
soilhorizon_thetas_10	[m <sup>3</sup> m <sup>-3</sup> ]	N(0.3160, 0.0316)
soilhorizon_thetar_10	[m <sup>3</sup> m <sup>-3</sup> ]	TN(0.0612, 0.0153, 0, 1)
soilhorizon_Ks_10	[m s <sup>-1</sup> ]	LN(5.91e-06, 5.5e-07)
soilhorizon_hg_10	[m]	N(-0.0356, 0.00356)
soilhorizon_mn_10	[-]	N(0.1791, 0.0179)
soilhorizon_Ko_10	[m s <sup>-1</sup> ]	LN(-6.24e-06, 5.5e-07)
soilhorizon_L_10	[-]	U(0.8376, 2.0940)
soilhorizon_bd_10	[g cm <sup>-3</sup> ]	U(1.2984, 1.9476)
soilhorizon_moc_10	[g g <sup>-1</sup> ]	U(0.0025, 0.0055)
soilhorizon_thetas_11	[m <sup>3</sup> m <sup>-3</sup> ]	N(0.3375, 0.0338)
soilhorizon_thetar_11	[m <sup>3</sup> m <sup>-3</sup> ]	TN(0.0372, 0.0093, 0, 1)
soilhorizon_Ks_11	[m s <sup>-1</sup> ]	LN(7.3e-06, 5.5e-07)
soilhorizon_hg_11	[m]	N(-0.0969, 0.00969)
soilhorizon_mn_11	[-]	N(0.2685, 0.0268)
soilhorizon_Ko_11	[m s <sup>-1</sup> ]	LN(-6.37e-06, 5.5e-07)
soilhorizon_L_11	[-]	U(-10.1124, -6.7416)
soilhorizon_bd_11	[g cm <sup>-3</sup> ]	U(1.0752, 1.6128)
soilhorizon_moc_11	[g g <sup>-1</sup> ]	U(0.0049, 0.0050)
soilhorizon_thetas_14	[m <sup>3</sup> m <sup>-3</sup> ]	N(0.3375, 0.0338)
soilhorizon_thetar_14	[m <sup>3</sup> m <sup>-3</sup> ]	TN(0.0372, 0.0093, 0, 1)
soilhorizon_Ks_14	[m s <sup>-1</sup> ]	LN(7.47e-06, 5.5e-07)
soilhorizon_hg_14	[m]	N(-0.0969, 0.00969)
soilhorizon_mn_14	[-]	N(0.2685, 0.0268)
soilhorizon_Ko_14	[m s <sup>-1</sup> ]	LN(-6.37e-06, 5.5e-07)
soilhorizon_L_14	[-]	U(-10.1124, -6.7416)
soilhorizon_bd_14	[g cm <sup>-3</sup> ]	U(1.0752, 1.6128)
soilhorizon_moc_14	[g g <sup>-1</sup> ]	U(0.0175, 0.0385)
soilhorizon_thetas_12	[m <sup>3</sup> m <sup>-3</sup> ]	N(0.3375, 0.0338)
soilhorizon_thetar_12	[m <sup>3</sup> m <sup>-3</sup> ]	TN(0.0372, 0.0093, 0, 1)
soilhorizon_Ks_12	[m s <sup>-1</sup> ]	LN(7.3e-06, 5.5e-07)
soilhorizon_hg_12	[m]	N(-0.0969, 0.00969)
soilhorizon_mn_12	[-]	N(0.2685, 0.0268)
soilhorizon_Ko_12	[m s <sup>-1</sup> ]	LN(-6.37e-06, 5.5e-07)
soilhorizon_L_12	[-]	U(-10.1124, -6.7416)
soilhorizon_bd_12	[g cm <sup>-3</sup> ]	U(1.0752, 1.6128)
soilhorizon_moc_12	[g g <sup>-1</sup> ]	U(0.0072, 0.0158)
soilhorizon_thetas_15	[m <sup>3</sup> m <sup>-3</sup> ]	N(0.3375, 0.0338)
soilhorizon_thetar_15	[m <sup>3</sup> m <sup>-3</sup> ]	TN(0.0372, 0.0093)
soilhorizon_Ks_15	[m s <sup>-1</sup> ]	LN(7.47e-06, 5.5e-07)
soilhorizon_hg_15	[m]	N(-0.0969, 0.00969)
soilhorizon_mn_15	[-]	N(0.2685, 0.0268)
soilhorizon_Ko_15	[m s <sup>-1</sup> ]	LN(-6.37e-06, 5.5e-07)
soilhorizon_L_15	[-]	U(-10.1124, -6.7416)
soilhorizon_bd_15	[g cm <sup>-3</sup> ]	U(1.0752, 1.6128)
soilhorizon_moc_15	[g g <sup>-1</sup> ]	U(0.0175, 0.0385)
soilhorizon_thetas_13	[m <sup>3</sup> m <sup>-3</sup> ]	N(0.3375, 0.0338)
soilhorizon_thetar_13	[m <sup>3</sup> m <sup>-3</sup> ]	TN(0.0372, 0.0093, 0, 1)
soilhorizon_Ks_13	[m s <sup>-1</sup> ]	LN(7.3e-06, 5.5e-07)
soilhorizon_hg_13	[m]	N(-0.0969, 0.00969)
soilhorizon_mn_13	[-]	N(0.2685, 0.0268)
soilhorizon_Ko_13	[m s <sup>-1</sup> ]	LN(-6.37e-06, 5.5e-07)
soilhorizon_L_13	[-]	U(-10.1124, -6.7416)
soilhorizon_bd_13	[g cm <sup>-3</sup> ]	U(1.0752, 1.6128)
soilhorizon_moc_13	[g g <sup>-1</sup> ]	U(0.0067, 0.0080)

Table B1. Continued.

Soil parameters		
soilhorizon_thetas_16	[m <sup>3</sup> m <sup>-3</sup> ]	N(0.3375, 0.0338)
soilhorizon_thetar_16	[m <sup>3</sup> m <sup>-3</sup> ]	TN(0.0372, 0.0093, 0, 1)
soilhorizon_Ks_16	[m s <sup>-1</sup> ]	LN(7.47e-06, 5.5e-07)
soilhorizon_hg_16	[m]	N(-0.0969, 0.00969)
soilhorizon_mn_16	[-]	N(0.2685, 0.0268)
soilhorizon_Ko_16	[m s <sup>-1</sup> ]	LN(-6.37e-06, 5.5e-07)
soilhorizon_L_16	[-]	U(-10.1124, -6.7416)
soilhorizon_bd_16	[g cm <sup>-3</sup> ]	U(1.0752, 1.6128)
soilhorizon_moc_16	[g g <sup>-1</sup> ]	U(0.0175, 0.0385)
Pesticide parameters		
pest_Koc	[m Lg <sup>-1</sup> ]	T(461.4000, 538.3000, 769.0000)
pest_DT50	[d]	N(47.1, 28.26)
Vegetation parameters		
veget_manning_1	[sm <sup>-1/3</sup> ]	T(0.0250, 0.0330, 0.041)
veget_Zr_1	[m]	U(2.096, 3.144)
veget_F10_1	[-]	U(0.2960, 0.4440)
veget_LAImin_1	[-]	U(0.0080, 0.0120)
veget_LAImax_1	[-]	U(2, 3)
veget_LAIharv_1	[-]	U(0.0080, 0.0120)
veget_manning_2	[sm <sup>-1/3</sup> ]	T(0.1000, 0.2000, 0.3000)
veget_Zr_2	[m]	U(7.2, 1.08)
veget_F10_2	[-]	U(0.2680, 0.4020)
veget_LAI_2	[-]	U(4, 6)
River parameters		
river_hpond	[m]	U(0.008, 0.012)
river_di	[m]	U(1.2, 1.8)
river_Ks	[m s <sup>-1</sup> ]	U(76.5648, 7.1280)
river_manning	[sm <sup>-1/3</sup> ]	T(0.0610, 0.0360, 0.0790)
Plot and VFS parameters		
plot_hpond	[m]	U(0.008, 0.012)
vfz_hpond	[m]	U(0.04, 0.06)
hu_adsorpthick	[m]	U(0.005, 0.015)

## Appendix C: Screening results

**Table C1.** Remaining parameters after the screening step for each output variable. In the XXX\_XXX\_XXX syntax of parameter names, the first block is the type of element the parameter refers to (soil horizon, river, vegetation, pesticide, HU or VFS), and the second part is the parameter name, while the last part is the element index the parameter refers to (soil horizon or vegetation type).

Water subsurface flow	Pesticide subsurface flow	Water surface runoff	Pesticide surface runoff
soilhorizon_thetas_2	soilhorizon_thetas_12	soilhorizon_thetas_11	soilhorizon_thetas_11
soilhorizon_thetas_4	soilhorizon_thetas_15	soilhorizon_thetas_15	soilhorizon_thetas_12
soilhorizon_thetas_6	soilhorizon_thetas_2	soilhorizon_thetas_2	soilhorizon_thetas_13
soilhorizon_thetas_7	soilhorizon_thetas_4	soilhorizon_thetas_4	soilhorizon_thetas_15
soilhorizon_thetas_8	soilhorizon_thetas_6	soilhorizon_thetas_6	soilhorizon_thetas_2
soilhorizon_thetas_10	soilhorizon_thetas_7	soilhorizon_thetas_7	soilhorizon_thetas_4
soilhorizon_thetar_3	soilhorizon_thetas_8	soilhorizon_thetas_8	soilhorizon_thetas_6
soilhorizon_thetar_4	soilhorizon_thetas_10	soilhorizon_thetas_10	soilhorizon_thetas_7
soilhorizon_thetar_8	soilhorizon_thetar_2	soilhorizon_thetar_15	soilhorizon_thetas_8
soilhorizon_thetar_10	soilhorizon_thetar_4	soilhorizon_thetar_2	soilhorizon_thetas_10
soilhorizon_moc_13	soilhorizon_thetar_8	soilhorizon_thetar_4	soilhorizon_thetar_15
soilhorizon_mn_3	soilhorizon_thetar_10	soilhorizon_thetar_8	soilhorizon_thetar_2
soilhorizon_mn_4	soilhorizon_pore_6	soilhorizon_thetar_10	soilhorizon_thetar_4
soilhorizon_mn_6	soilhorizon_moc_12	soilhorizon_pore_9	soilhorizon_thetar_8
soilhorizon_mn_8	soilhorizon_moc_15	soilhorizon_mn_11	soilhorizon_thetar_10
soilhorizon_mn_10	soilhorizon_moc_2	soilhorizon_mn_2	soilhorizon_moc_6
soilhorizon_Kx_3	soilhorizon_moc_6	soilhorizon_mn_4	soilhorizon_moc_12
soilhorizon_Kx_4	soilhorizon_moc_9	soilhorizon_mn_6	soilhorizon_mn_11
soilhorizon_Kx_8	soilhorizon_mn_11	soilhorizon_mn_7	soilhorizon_mn_16
soilhorizon_Kx_10	soilhorizon_mn_16	soilhorizon_mn_8	soilhorizon_mn_2
soilhorizon_Ks_11	soilhorizon_mn_4	soilhorizon_mn_10	soilhorizon_mn_4
soilhorizon_Ks_13	soilhorizon_mn_6	soilhorizon_Kx_8	soilhorizon_mn_6
soilhorizon_Ks_14	soilhorizon_mn_8	soilhorizon_Kx_10	soilhorizon_mn_7
soilhorizon_Ks_15	soilhorizon_mn_10	soilhorizon_Ks_12	soilhorizon_mn_8
soilhorizon_Ks_16	soilhorizon_Kx_12	soilhorizon_Ks_13	soilhorizon_mn_10
soilhorizon_Ks_3	soilhorizon_Kx_9	soilhorizon_Ks_15	soilhorizon_Ks_15
soilhorizon_Ks_4	soilhorizon_Kx_10	soilhorizon_Ks_16	soilhorizon_Ks_2
soilhorizon_Ks_6	soilhorizon_Ks_12	soilhorizon_Ks_4	soilhorizon_Ks_4
soilhorizon_Ks_7	soilhorizon_Ks_14	soilhorizon_Ks_6	soilhorizon_Ks_8
soilhorizon_Ks_8	soilhorizon_Ks_15	soilhorizon_Ks_8	soilhorizon_Ks_9
soilhorizon_Ks_9	soilhorizon_Ks_16	soilhorizon_Ks_9	soilhorizon_hg_3
soilhorizon_Ks_10	soilhorizon_Ks_2	soilhorizon_Ks_10	soilhorizon_hg_4
soilhorizon_hg_16	soilhorizon_Ks_4	soilhorizon_hg_4	soilhorizon_hg_8
soilhorizon_hg_2	soilhorizon_Ks_6	soilhorizon_hg_6	soilhorizon_bd_6
soilhorizon_hg_4	soilhorizon_Ks_8	soilhorizon_hg_8	soilhorizon_bd_13
soilhorizon_hg_8	soilhorizon_Ks_9	soilhorizon_hg_10	soilhorizon_bd_12
soilhorizon_hg_9	soilhorizon_Ks_10	soilhorizon_bd_3	river_ks
soilhorizon_bd_2	soilhorizon_hg_4	river_ks	river_di
river_ks	soilhorizon_hg_6	river_di	plot_hpond
river_di	soilhorizon_hg_8	plot_hpond	pest_Koc_1
plot_hpond	soilhorizon_hg_10	vfz_hpond	pest_DT50_1
VFS_hpond	soilhorizon_bd_11	veget_LAlharv_1	HU_adsorpthick
	soilhorizon_bd_12	veget_F10_1	veget_Zr_1
	soilhorizon_bd_13		veget_manning_1
	soilhorizon_bd_15		veget_F10_1
	soilhorizon_bd_2		
	soilhorizon_bd_6		
	soilhorizon_bd_9		
	river_ks		
	river_di		
	plot_hpond		
	veget_Zr_1		
	pest_Koc_1		
	pest_DT50_1		

*Code and data availability.* The PESHMELBA model is an open-source model coded in Python (Version 2.7.17) and Fortran 90 and embedded in the OpenPALM coupler (Version 4.3.0). The code for the OpenPALM coupler is available from [http://www.cerfacs.fr/globc/PALM\\_WEB/user.html#download](http://www.cerfacs.fr/globc/PALM_WEB/user.html#download) (Fouilloux and Piacentini, 1999) after registration. The exact version of PESHMELBA used to produce the simulations is archived on Zenodo (<https://doi.org/10.15454/2HAU8R>, Rouzies et al., 2022a) as are input data and scripts used to produce all the sensitivity indices presented in this paper (<https://doi.org/10.15454/2YVY4O>, Rouzies et al., 2022b). These scripts use the open-source MATLAB software for uncertainty quantification UQLab Version 2.0.0 (<https://www.uqlab.com>, Marelli and Sudret, 2014). The R package randomForestSRC is freely available for download at <https://cran.r-project.org> (R Core Team, 2017).

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