Causal Inference in Geoscience and Remote Sensing from Observational Data

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Abstract—Establishing causal relations between random variables from observational data is perhaps the most important challenge in today’s science. In remote sensing and geosciences this is of special relevance to better understand the Earth’s system and the complex interactions between the governing processes. In this paper, we focus on observational causal inference, thus we try to estimate the correct direction of causation using a finite set of empirical data. In addition, we focus on the more complex bivariate scenario that requires strong assumptions and no conditional independence tests can be used. In particular, we explore the framework of (non-deterministic) additive noise models, which relies on the principle of independence between the cause and the generating mechanism. A practical algorithmic instantiation of such principle only requires 1) two regression models in the forward and backward directions, and 2) the estimation of statistical independence between the obtained residuals and the observations. The direction leading to more independent residuals is decided to be the cause. We instead propose a criterion that uses the sensitivity (derivative) of the dependence estimator, the sensitivity criterion allows to identify samples most affecting the dependence measure, and hence the criterion is robust to spurious detections. We illustrate performance in a collection of 28 geoscience causal inference problems, in a database of radiative transfer models simulations and machine learning emulators in vegetation parameter modeling involving 182 problems, and in assessing the impact of different regression models in a carbon cycle problem. The criterion achieves state-of-the-art detection rates in all cases, it is generally robust to noise sources and distortions. The presented approach confirms the validity in observational bi-variate problems in the Earth sciences.

Index Terms—Causal inference, Dependence estimation, Regression, Noise, Sensitivity, Hilbert-Schmidt Independence Criterion (HSIC), Gaussian Process

I. INTRODUCTION

"... observational studies are an interesting and challenging field which demands a good deal of humility, since we can claim only to be groping toward the truth.”

William Cochran (1972) [1].

THE Earth is a highly complex and evolving networked system that we strive to understand better to deal with societal, economical and environmental challenges, such as climate change [2], [3]. There is an urgent need for tools that help us observe and study the Earth system. Nowadays, machine learning and signal processing play a crucial role for the production and analysis of Earth observation empirical data provided by a plethora of sensory systems and platforms. However, most statistical methods focus on prediction and estimation problems so they are only designed to take advantage of association relationships without considering causal mechanisms. Such methods provide little information about how variables actually interact with each other and, in this sense, are not very helpful to understand the underlying processes governing the system. The purpose of causal inference is precisely to go beyond association and to determine and discover links of causes and effects. Unlike association (e.g. correlation) studies, causal studies allow for understanding the underlying processes, and thus enable making inferences (i.e. predictions) of the effects of actions on the observed system [4]–[6].

Causal inference should be ideally performed through the design of controlled experiments that try to avoid variable selection and confounding biases. Considering all possible variables and controlling all possible interactions would be of course the ideal scenario. Setting up such experiments is, however, not always possible, notably for ethical, economical or simply feasibility reasons [7]. This is often the case in empirical sciences, such as remote sensing, climate science and the geosciences, where one cannot control the whole set of variables affecting a given experiment. This is why actual causal experiments on the Earth system are often replaced by factorial experiments done with an ensemble of Earth-system model simulations [8]. Actually, a vast literature collectively perform model-based causal inference, and they focus on climate data only. The studies typically rely on climate models [9], and explore schemes for detection and attribution of plausible causes by running models under different scenarios that consider or not the variable (forcing) under inspection.

The main advantages of model-based approaches is that, in general, models encapsulate all the current knowledge about the system and thus they account for all the factors that can influence it. However, even though climate, biogeochemical or radiative transfer models are based on well-known physics equations, the combination of all internal processes and their couplings still make the interpretation of outputs very complicated, and it turns to be very difficult to disentangle internal from external induced variabilities. It is also worth noting that models are not a perfect representation of reality and many assumptions are made, so the eventual conclusions derived from these studies can be limited, or even wrong.
As an alternative to the use of models, one can resort to pure observational data, which opens a wide field in the current era of data deluge. It is acknowledged that the problem of inferring causation from empirical data has been traditionally considered unsolvable. Indeed, given two variables, identifying which is the cause and which one is the effect requires adopting (strong) assumptions. For example, it is assumed the absence of ‘confounding factors’ that might drive both variables. This means that the system is fully described by two variables only, so there is not a third variable driving both. A second important assumption is known as the problem of ‘selection bias’. This implies that the observed variables should be representative of the causal relationship. A final third assumption commonly adopted is that no feedback loops can be found or created neither [4]. A plethora of methods of causal discovery exist that try to remedy such limitations by 1) considering all potentially explanatory variables of the phenomenon, 2) selecting the most impactful ones, and 3) estimating conditional independence between (subsets of) observed variables to create directed graphs. Advances in observational causal inference have permitted to draw (partial) conclusions about the causal relationships in real-life problems [4], [5], [10], [11]. In remote sensing and geosciences, this is of special relevance to better understand the Earth’s system and the complex and elusive interactions between the involved processes. Answering key questions may have deep societal, economical and environmental implications [3], [12].

In the field of remote sensing and geosciences, two main methodological approaches exist, and both of them consider the multivariate setting and that time is involved. Time gives an obvious intuition on causality (“the cause should precede the effect”) and having access to multiple time series allows to overcome the selection bias problem. The seminal work by Granger [13] proposed a rather simple statistical concept of causality based on prediction, and has been applied in many fields of Earth system science: to perform attribution of climate change [9], [14]–[16]; to study the feedback mechanism between soil moisture and precipitation [17]; or to identify the relationship between sea surface temperature (SST) and the North Atlantic Oscillation (NAO) [18]. Recently, Granger causality has been adapted to consider nonlinear relations in vegetation dynamics [19]. As an alternative to prediction, a second family of methods consider that it is sometimes possible to retrieve, at least partially, the causal structure using conditional independence tests between the variables in PC schemes [20]. The family is called constraint-based search, and has been used to study the causal interactions between climate modes of variability [21], as well as to construct climate causal networks [22], [23]. Similar methodologies were applied to study the causal relationships with the Atlantic meridional overturning circulation (AMOC) [24], to analyze potential drivers of Artic Oscillation [25], or to investigate the interactions between El Niño-Southern Oscillation (ENSO) and the Walker circulation [26].

In this paper, unlike in the previous approaches, we will focus on the more challenging problem of inferring causality from observational data with two important constraints:

- **Bivariate case.** We will consider the case of having access to two variables only, hence no conditional independence tests can be computed to guide the causal identification.
- **Time is not explicitly considered.** We will not consider time-series explicitly in general, hence we deal with the problem of ‘instantaneous causality’. This hampers the adoption of the ‘causes precede effects’ rationale, so specific time-series approaches such as those based on Granger causality methods cannot be applied. Note that, however, time is often not necessary to discuss concepts such as statistical dependence and, in causal models, time is often not even needed to discuss the effect of interventions.

We call this setting the bivariate instantaneous case, which has been recently treated in [6], [27]. Let us thus assume that only two variables, \(x\) and \(y\), have been observed, and we aim to distinguish \(x\) causing \(y\) (indicated as \(x \rightarrow y\)) from \(y\) causing \(x\) (that is \(y \rightarrow x\)) using only purely observational data, i.e., a finite i.i.d. sample drawn from the joint distribution \(p(x,y)\). Following the Bayes’ theorem, there are two admissible partitions of the joint distribution:

\[
p(x,y) = p(y|x)p(x) = p(x|y)p(y),
\]

and the question is to decide which one is the causal one. The first one describes variable \(x\) and the conditional \(p(y|x)\) that can be interpreted as a function that translates the information contained in \(x\) to \(y\), so it assumes that \(x\) causes \(y\). The second decomposition assumes that modeling \(y\) and the conditional \(p(x|y)\) better explains the joint distribution, thus assuming that \(y\) causes \(x\). A possible answer to this question could be obtained through an intervention analysis (alter a variable and see the plausibility of the effect on the system), but this is not possible in our Earth science problems for obvious reasons. An alternative pathway is to rely on the principle of independent mechanisms [6], [28]: if we assume \(x \rightarrow y\) so the causal partition is \(p(x,y) = p(y|x)p(x)\), one would expect the conditional density \(p(y|x)\) to provide no information about the marginal density \(p(x)\), or at least less information than \(p(x|y)\) would provide about \(p(y)\). Therefore, a solution to the bivariate case reduces to estimate independence between the cause density and the mechanism producing the effect distribution. This is called the ‘independence of cause and mechanism’ (ICM) [29], [30], which is the one adopted in this paper. We should note that not all systems satisfy this principle. The problem is to achieve good density estimates of the conditionals and potential causes, and to measure independence between them accurately.

Following the previous rationale, and focusing on the non-deterministic scenario, we here explore the framework of nonlinear additive noise models (ANMs) [31], which rely on the principle of independence between the cause (prior) and the generating mechanism (conditional). A practical algorithmic instantiation of such principle only requires 1) two regression models to learn the two possible forward and backward directions of causation, and 2) the estimation of statistical independence between the obtained residuals by the factorization and the observation. The direction leading to more independent residuals is decided to be the cause. The method has achieved...


state-of-the-art results in an exhaustive comparison involving bivariate causal problems in biology, geosciences, economy and social sciences, as illustrated in [27]. Authors in [31] suggested the use of Gaussian processes [32], [33] for regression fitting arbitrarily complex functions, and the Hilbert Schmidt Independence Criterion (HSIC) [34] as dependence test based on the excellent convergence properties to the true dependence and ease of calculation. HSIC has been previously used in remote sensing for feature selection and dependence estimation [35], [36].

We here propose an alternative criterion to the direct dependence of the residuals, and focus on the sensitivity (derivative) of the HSIC dependence estimate. The paper extends our previous work in [37] with more theoretical insight and a larger set of experimental evidence. In particular, we illustrate performance in a collection of 28 geoscience causal inference problems, in a large database of radiative transfer models simulations and machine learning emulators in vegetation parameter modeling leading to 182 causal problems with ground truth, and in assessing the impact of different regression models in a carbon cycle problem. The criterion achieves state-of-the-art detection rates above chance levels, it is robust to noise sources and distortions, and the adoption of different regression models. The presented approach confirms the validity in observational bi-variate problems in the Earth sciences.

The remainder of the paper is organized as follows. Section II reviews the main aspects of the adopted causal framework, and the needed tools for the practical implementation: Gaussian processes for regression and the HSIC estimate for dependence estimation. Section III derives the HSIC sensitivity maps and describes the proposed causal criterion. Section IV gives experimental evidence of performance in a wide range of bivariate Earth system problems. Finally, we conclude in Section V with some remarks and future outlook.

II. CAUSALITY IN PAIRS OF INSTANTANEOUS VARIABLES

A. Deterministic setting

Let us first consider the case where noise is not present. This deterministic causal problem has been treated before [6], [38], [39], and is often known as Information-Geometric Causal Inference (IGCI). Formally, let us assume a continuous differentiable transformation, \( f : x \to y \). The function \( f \) has to be a diffeomorphism (it is differentiable and bijective and it has a differentiable inverse) of \([0, 1]\) that is strictly monotonic and satisfies \( f(0) = 0 \) and \( f(1) = 1 \). Now, using the standard formula of densities under transformation

\[
p(x) = p(y) \left| \frac{\partial f(x)}{\partial x} \right|,
\]

one could arguably identify the direction of causation from the particular structure of the densities. If the structure of \( p(x) \) is not correlated with the slope of \( f \), then flat regions of \( f \) induce peaks in \( p(y) \). The causal hypothesis \( y \to x \) is thus implausible because the causal mechanism \( f^{-1} \) appears to be adjusted to the ‘input’ distribution \( p(y) \). Note that here the principle of independence of cause and mechanism reduces to estimate the independence of \( p(x) \) and \( f \), which interestingly implies dependence between \( p(y) \) and \( f^{-1} \). This is illustrated in Fig. 1.

We should note, however, that such justifications always refer to oversimplified models that are unlikely to describe realistic situations. After all, bijective deterministic relations are rare in nature. Therefore, IGCI only provides a limited, unrealistic scenario for which cause-effect inference is possible by virtue of an approximate cause-mechanism independence assumption.

B. Non-deterministic setting

The vast majority of real (and thus more interesting) problems are not deterministic, the function \( f \) relating the two variables is not bijective, and \( f \) is inaccessible so one cannot compute its derivatives, and from there a criterion of independence between the densities of the cause and the mechanism. An alternative is to rely on the assumption made in Additive Noise Models (ANM) for causal inference [31].

Given two random variables \( x \) and \( y \) with causal relation \( x \to y \), under some conditions, one can demonstrate (cf. e.g. Theorem 4.5 about the identifiability of ANMs in [6]) that there exists an additive noise model

\[
y = f(x) + n_f,
\]

in the correct causal direction, but there exists no additive noise model

\[
x = g(y) + n_g
\]
in the anticausal direction. The above mentioned Theorem can be derived through the particular assumptions and it states that generically, a distribution does not admit an ANM in both directions at the same time.

This observation allowed Hoyer et al. [31] to design an algorithm to distinguish cause from effect in pairs of variables from empirical data. Essentially, the methodology performs nonlinear regression from $x \to y$ (and vice versa, $y \to x$) and assesses the independence of the forward, $r_f = y - f(x)$, and backward residuals, $r_b = x - g(y)$, with the input variables (drivers) $y$ and $x$, respectively. The more independent residuals tell us the right direction of causation.

Relevant assumptions are done in this approach. First, we assume that the considered problem is described accurately looking at the pairs (representational property). Another important assumption is the causal sufficiency, which states that there is no hidden common cause in the considered variables that is causing any of the latter, and thus acts as a confounder. Third, the causal Markov assumption holds, which allows us to use the causal graph as a probabilistic one. In addition, there are two extra conditions on the regression functions: (i) $f, g$ are either linear and noise is non-Gaussian, or (ii) $f, g$ are nonlinear and output’s densities are positive and smooth [6], [27].

In order to define a practical criterion, two ingredients are needed only: a regression method to learn functions $f$ and $g$, and a powerful dependence estimate to assess the independence of the residuals. Therefore, the framework needs two fundamental blocks: 1) a nonlinear regression model, and 2) a dependence measure. We typically rely on Gaussian Processes [33] and the HSIC [34], respectively. In what follows, we briefly review the theory under these two kernel methods [40]. Then we define the causality criterion.

1) Gaussian Processes (GPs): Standard regression approximates observations (often referred to as outputs) as the sum of some unknown latent function $f(x)$ of the input data plus constant power (homoscedastic) Gaussian noise of variance $\lambda^2$. Note that in our case, both inputs and outputs are unidimensional, $x, y \in \mathbb{R}$. Therefore, given $n$ input-output data pairs, the dataset is denoted as $\mathcal{D} = \{(x_i, y_i)\}_{i=1}^n$, and the model approximation is

$$y_i = f(x_i) + \varepsilon_i, \quad \varepsilon_i \sim \mathcal{N}(0, \lambda^2).$$

Instead of proposing a parametric form for $f(x)$ and learning its parameters in order to fit observed data well, GP regression proceeds in a Bayesian, non-parametric way. A zero mean GP prior is placed on the latent function $f(x)$ and a Gaussian prior is used for each latent noise term $\varepsilon_i$, $f(x) \sim \mathcal{GP}(0, k_\theta(x, x'))$, where $k_\theta(x, x')$ is a covariance function parametrized by $\theta$ and $\sigma^2$ is a hyperparameter that specifies the noise power. Essentially, a GP is a stochastic process whose marginals are distributed as a multivariate Gaussian. In particular, given the priors $\mathcal{GP}$, samples drawn from $f(x)$ at the set of locations $\{x_i\}_{i=1}^n$ follow a joint multivariate Gaussian with zero mean and covariance matrix $K_{ff}$ with $[K_{ff}]_{ij} = k_\theta(x_i, x_j)$.

If we consider a test location $x_s$ with corresponding output $y_s$, priors $\mathcal{GP}$ induce a prior distribution between the observations $y = [y_1, \ldots, y_n]^\top$ and $y_s$. Now it is possible to analytically compute the posterior distribution over the unknown output $y_s$ given the test input $x_s$ and the available training set $\mathcal{D}$,

$$p(y_s|x_s, \mathcal{D}) = \mathcal{N}(y_s | \mu_{\mathcal{GP}}, \sigma_{\mathcal{GP}}^2),$$

which is a Gaussian with the following mean and variance:

$$\mu_{\mathcal{GP}} = k_f^\top (K_{ff} + \lambda^2I_n)^{-1} y$$

$$\sigma_{\mathcal{GP}}^2 = \lambda^2 + k_{ss} - k_f^\top (K_{ff} + \lambda^2I_n)^{-1} k_f,$$

where $k_f = [k(x_s, x_1), \ldots, k(x_s, x_n)]^\top \in \mathbb{R}^n$ contains the kernel similarities of the test point $x_s$ to all training points in $\mathcal{D}$, $K_{ff}$ is a $n \times n$ kernel (covariance) matrix whose entries contain the similarities between all training points, $\lambda^2$ is a hyperparameter accounting for the variance of the noise, $k_{ss} = k(x_s, x_s)$ is a scalar with the self-similarity of $x_s$, and $I_n$ is the identity matrix of size $n$. Note that both the predictive mean and the variance can be computed in closed-form, and that the predictive variance $\sigma_{\mathcal{GP}}^2$ does not depend on the outputs/target variable.

2) Kernel Dependence Estimation with HSIC: Let us consider two spaces $\mathcal{X}, \mathcal{Y} \subseteq \mathbb{R}$, on which we jointly sample observation pairs $(x, y)$ from distribution $p(x, y)$. The covariance matrix $C_{xy}$ encodes first order dependencies between the random variables. A statistic that efficiently summarizes the content of this matrix is its Hilbert-Schmidt norm. This quantity is zero if and only if there exists no second order dependence between $x$ and $y$.

The nonlinear extension of the notion of covariance was proposed in [34] to cope with higher-order relations between the data. The use of its linear formulation has some limitations and cannot discover more complex relations, for this purpose the use of nonlinear kernel functions allows to capture higher-order effects. Let us define a (possibly nonlinear) mapping $\phi : \mathcal{X} \to \mathcal{F}$ such that the inner product between features is given by a positive definite (p.d.) kernel function $K_x(x, x') = \langle \phi(x), \phi(x') \rangle$. The feature space $\mathcal{F}$ has the structure of a reproducing kernel Hilbert space (RKHS). Let us now denote another feature map $\psi : \mathcal{Y} \to \mathcal{G}$ with associated p.d. kernel function $K_y(y, y') = \langle \psi(y), \psi(y') \rangle$. Then, the cross-covariance operator between these feature maps is a linear operator $C_{xy} : \mathcal{G} \to \mathcal{F}$ such that $C_{xy} = \mathbb{E}_{xy}[(\phi(x) - \mu_x) \otimes (\psi(y) - \mu_y)]$, where $\otimes$ is the tensor product, $\mu_x = \mathbb{E}_x[\phi(x)]$, and $\mu_y = \mathbb{E}_y[\psi(y)]$. See more details in [41], [42]. The squared norm of the cross-covariance operator, $||C_{xy}||_{HS}^2$, is called the Hilbert-Schmidt Independence Criterion (HSIC) and can be expressed in terms of kernels [34]. Given the set $\mathcal{D}$ with $n$ scalar pairs drawn from the joint $p(x, y)$ an empirical estimator of HSIC is [43]:

$$\text{HSIC} = \frac{1}{n} \text{Tr}(HK_xH K_y),$$
The formulation of the HSIC sensitivity maps, the proposed causality criterion, and study their empirical and theoretical properties.

A. Sensitivity analysis and maps

A general definition of the sensitivity map (SM) in the context of kernel methods was originally introduced in [45]. Let us define HSIC as a function \( h : \mathcal{X} \times \mathcal{Y} \to \mathbb{R} \). The sensitivity map is the expected value of the squared derivative of the function (or the log of the function) with respect to the arguments \( z = (x, y) \). Formally, let us define the sensitivity as

\[
\begin{align*}
\text{s} &= \int_{\mathcal{Z}} \left( \frac{\partial h(z)}{\partial z} \right)^2 p(z) dz, \\
&= \int_{\mathcal{Z}} \left( \frac{\partial h(z)}{\partial z} \right)^2 dz,
\end{align*}
\]

(7)

where \( p(z) \) is the probability density function over the inputs \( z \in \mathcal{Z} \). Intuitively, the objective of the sensitivity map is to measure the changes of the function \( h(x) \) along the inputs \( x \). In order to avoid the possibility of cancellation of the terms due to its signs, the derivatives are typically squared, even though other unambiguous transformations like the absolute value could be equally applied. Integration gives an overall measure of sensitivity over the observation space \( \mathcal{Z} \). The empirical sensitivity map approximation to Eq. (7) is obtained by replacing the integral with a summation over the available \( n \) samples

\[
\begin{align*}
\text{s} &\approx \frac{1}{n} \sum_{i=1}^{n} \left( \frac{\partial h(x_i)}{\partial z} \right)^2.
\end{align*}
\]

(8)

The intuition behind the approach is that statistically significant residuals in one direction indicates the true data-generating mechanism (see Sections §I and §II-A). See an illustrative example in Fig. 2. The problem contains values of generating mechanism (see Sections §I and §II-A). See an illustrative example in Fig. 2. The problem contains values of generating mechanism (see Sections §I and §II-A). See an illustrative example in Fig. 2. The problem contains values of generating mechanism (see Sections §I and §II-A). See an illustrative example in Fig. 2. The problem contains values of generating mechanism (see Sections §I and §II-A). See an illustrative example in Fig. 2. The problem contains values of generating mechanism (see Sections §I and §II-A). See an illustrative example in Fig. 2. The problem contains values of generating mechanism (see Sections §I and §II-A). See an illustrative example in Fig. 2. The problem contains values of generating mechanism (see Sections §I and §II-A).
Since the HSIC function $h$ works on $(x, y)$, we have to apply this expression twice, which returns the sensitivity vector $s = [s^x, s^y]$. This gives the relevance of variables $x$ and $y$ in the function $h$. Alternatively, one can summarize the maps variable-wise to obtain a point-wise relevance:

$$p_i = \sqrt{\left(\frac{\partial h(x)}{\partial x}\right)^2 + \left(\frac{\partial h(y)}{\partial y}\right)^2},$$

(9)

which will be used in this paper to evaluate the relevance of points in the dependence measure. Actually, this way of summarizing the information conveyed by the sensitivity map is somewhat related to the concepts of leveraging and influential points in statistics [46].

B. Sensitivity maps for the HSIC

In order to derive the sensitivity maps for HSIC which were originally presented in [47], we need to compute its partial derivatives w.r.t. points in variables $x$ and $y$, i.e. $x_i$ and $y_i$. By applying the chain rule, and first-order derivatives of matrices, we obtain:

$$s^x_i := \frac{\partial \text{HSIC}}{\partial x_i} = -\frac{2}{\sigma^2 n^2} \text{Tr}(HK_xH(K_x \circ M_i)), \quad (10)$$

where matrix $M_i$ depends on the $i$-th point sample and it is formed by zeroes except in the $i$-th column which corresponds to vec$(x_i) - x$ ($i$-th element minus vector $x$).

$$s^y_i := \frac{\partial \text{HSIC}}{\partial y_i} = -\frac{2}{\sigma^2 n^2} \text{Tr}(HK_yH(K_y \circ N_i)), \quad (11)$$

where matrix $N_i$ depends on the $i$-th point sample and it is formed by zeroes except in the $i$-th column which corresponds to vec$(y_i) - y$ ($i$-th element minus vector $y$).

The joint sensitivity map is defined as the concatenation of individual sensitivities, $s = [s^x, s^y]$, where $s^x = \frac{1}{n} \sum_{i=1}^n (s^x_i)^2$ and $s^y = \frac{1}{n} \sum_{i=1}^n (s^y_i)^2$; and the point-wise sensitivity as $p_i = \sqrt{(s^x_i)^2 + (s^y_i)^2}$.

C. Proposed causal criterion with sensitivity maps

We here propose an alternative criterion for bivariate causality based on the sensitivity maps of HSIC in both directions:

$$C_s := (s^b_f + s^f_b) - (s^f_f + s^f_f), \quad (12)$$

where subscripts $f$ and $b$ stand for the forward and backward directions respectively, and the superscripts refer to the sensitivities of either the observations $x$ and $y$, or the corresponding residuals, $r_f$ and $r_b$. The criterion now accounts for the relative relevance of points and residuals in the dependence estimate according to the sensitivity measure. Besides, note the intuitive connection to the deterministic case in section II-A. Here we do not focus on the derivative of the underlying function, but the derivative of the dependence estimate itself [39]. Following the same “altitude causes temperature” example, we show in Fig. 3 the point sensitivities in the distributions. It becomes clearer the structured (more dependent) distribution in the backward direction (Fig. 3[right]), which suggests that the forward direction is the causal one.

D. Consistency Properties

Let us now study the consistency properties of proposed criterion. Following [44], a plausible criterion of causality for ANMs has to rely on a consistent dependence criterion, and the combination of the ANM and the dependence estimator should be consistent too.

Let us show first that the norm of the sensitivity map acts as a consistent dependence measure. Note that in [31] originally proposed to use the $p$-value of HSIC to define the causality criterion which leads to a consistent procedure, and in [44] it is shown that the same happens for the HSIC itself. Therefore, we have to ensure here that the norm of the sensitivity maps acts as a consistent dependence measure. It is customary to consider that if an estimator $\rho(X, Y)$ fulfills that “$\rho(X,Y) = 0$ iff $X$ and $Y$ are statistically independent”, then it is considered a dependence measure. Let us first give an empirical demonstration that the norm of the sensitivity maps of HSIC is a consistent estimator: under the hypothesis of independence (Fig. 4[a-b]) both HSIC and $\|\text{HSIC}\|$ converge to zero with $N$; while in an example of increasing dependence (Fig. 4[c]) convergence is ensured too.

In theoretical terms, to ensure consistency of the proposed criterion, we must demonstrate that for each of the four terms in $\tilde{C}_s := C_b - C_f = (s^b_f + s^f_b) - (s^f_f + s^f_f)$.

We will show it for the forward term $C_f$, the same arguments apply for $C_b$:

$$C_f(x, r) := s^y_f + s^r_f = \|\partial_x \text{HSIC}\|^2 + \|\partial_y \text{HSIC}\|^2.$$

Note that for real numbers, the norm of the sensitivity maps of HSIC should be Lipschitz continuous to ensure convergence, that is, we need to demonstrate $|C_f(x, r) - C_f(x, r')| \leq L \|r - r'\|$. For that, we can show that a multivariate function $f : \mathbb{R}^D \to \mathbb{R}$ with bounded partial derivatives, which is our case, is Lipschitz, and that the bound is exactly $L = \sqrt{2} \max_i (\sup \|\partial_i \text{HSIC}\|)$, $i = 1, \ldots, N$. This is an equivalent result to Lemma 16 in [44] for the HSIC sensitivity map.

The second condition to fulfill is that the norm of the sensitivity map in ANMs should be consistent. This can be demonstrated following the same procedure that the one in Theorem 20 (Appendix A.3) in [44] for HSIC, and using the previous consistency lemma for the proposed estimator here.
E. Computational cost and efficient criterion

Note that both HSIC and its sensitivity map give raise to closed-form solutions, just involving simple matrix multiplication and a trace operation. Both HSIC and its sensitivity scale quadratically with the number of examples $N$ since the involved kernel matrices and the centering matrix are $N \times N$. This makes both HSIC and its sensitivity map unfeasible with moderate to large datasets. In [47] we provided with fast versions of both HSIC and its sensitivity map through the use of random Fourier features. The cost of a naive implementation of HSIC is $O(N^3)$, and its randomized version is $O(D^3)$, where $D$ is the number of Fourier features chosen to approximate the kernel matrices, which is typically smaller than the number of points, i.e. $D \ll N$. Respectively, the naive implementation of the sensitivity of HSIC is $O(N^3 + N^2) \approx O(N^3)$ and its randomized version scales as $O(D^3 + D^2 N)$ so in the cases of $N \gg D$ the cost reduces to $O(D^2 N)$. The cost of the causal criteria is thus defined by these operations. In addition, we demonstrated the convergence of both the randomized HSIC to HSIC, and their corresponding sensitivity maps, which allows a practical use of the method.

IV. RESULTS

In this section we show the performance of the proposed methodology in three experimental settings: (1) in a collection of 28 geoscience causal inference problems, (2) in a database of radiative transfer models simulations and machine learning emulators in Sentinel-2 vegetation parameter modeling conforming a set of 182 causal problems with groundtruth, and (3) in assessing the impact of different regression models based on GPs in discovering causality in Net Ecosystem Production variables. All problems are bivariate and a annotated ground truth is available based on expert knowledge or common sense, which allows us to assess performance quantitatively.

We quantify the accuracy in detecting the direction of causation using standard scores like the receiver operating curves (ROC), precision-recall curves (PRC) curves, and the areas under these curves. We compare in all cases the state-of-the-art criterion in (6) with our criterion in (12). Methods performance are also studied under different situations of number of available points, presence of different noise sources and distortions, and impact of different GP regression models on the detection accuracy.
A. Experiment 1: Geoscience Cause Effect Pairs

We used Version 1.0 of the CauseEffectPairs (CEP) collection freely\(^2\). The database contains 100 pairs of random variables along with the right direction of causation (ground truth). Data has been collected from various domains of application, such as biology, climate science, health sciences and economics, just to name a few. More information about the dataset and an excellent up-to-date review of observational causal inference methods is available in [27]. We conducted experiments in 28 out of the 100 pairs that contain one-dimensional variables and that are related to geosciences and remote sensing: problems involving carbon and energy fluxes, ecological indicators, vegetation indices, temperature, moisture, heat, etc. We summarize the involved variables in Table I, and show the scatter plots of the selected pairs \((x, y)\) in Fig. 5.

1) Experimental Setup: The experimental setting is as follows. Once the two predictive models \(f\) and \(g\) were developed, we computed the two HSIC terms, as well as their sensitivity maps between the \(x\) (or \(y\)) and the residuals \(r_f\) (or \(r_b\)). The final causal direction score was simply defined as the difference in test statistic between both models, either using \(\hat{C}\) or the proposed \(\hat{C}_s\). Note that this is a particular form of ‘ranked-decision’ setting that needs to account for the bias introduced by pairs coming from the same problem, i.e. it is customary to down-weight the precision for every decision threshold in the curves (e.g. four related problems receive 0.25 weights in the decision)\(^3\). This is the case, for example of problems 81, 82, 83 and 87 that receive 0.25 weights.

2) Accuracy and robustness of the detection: We run the experiments with different numbers of (randomly selected) points \(n\) from both variables. This situation impacts regression models performance, both in terms of the regression accuracy and the dependence estimation. We evaluate \(\hat{C}\) and \(\hat{C}_s\) criteria by limiting the maximum number of training samples in each problem, \(n_{\text{max}} = \{50, 100, 200, 500, 2000\}\). Results were averaged over 10 realizations. Figure 6 shows the AUC under the curve as a function of \(n_{\text{max}}\). The proposed sensitivity-based criterion consistently performed better than the standard approach using HSIC alone. Lower values of AUC are obtained by our proposal only when the number of samples is relatively low (50 and 100). This particular behavior is plausible because our criterion is defined through an empirical estimator (the sensitivity) and when the number of samples is moderately low it can give rise to underestimated values. Looking more in detail at the ROC in Fig. 7, we note that the proposed \(\hat{C}_s\) yields the best recognition curves, both ROC and PRC. Note that this happens for all decision rates, especially in low false positive rates regimes, and for all number of training points.

\(^2\)https://webdav.tuebingen.mpg.de/cause-effect/
\(^3\)The MATLAB function perfcurve can produce such (weighted) ROC and PRC curves and the estimated weighted AUC.

B. Experiment 2: Causation in RTM assessment

Using input-output data pairs generated by radiative transfer models (RTMs) allow us to assess performance of observational causality algorithms: it is obvious that the forward
RTM simulation gives the right direction of causation: state vectors (parameters) cause radiances and not the other way around. The strength of the causation is not considered here. In this second experiment, we deal with data generated by the standard PROSAIL RTM. PROSAIL is the combination of the PROSPECT leaf optical properties model and the SAIL canopy bidirectional reflectance model. PROSAIL has been used to develop new methods for retrieval of vegetation biophysical properties. Essentially, PROSAIL links the spectral variation of canopy reflectance, which is mainly related to leaf biochemical contents, with its directional variation, which is primarily related to canopy architecture and soil/vegetation contrast. This link is key to simultaneous estimation of canopy biophysical/structural variables for applications in agriculture, plant physiology, and ecology at different scales. PROSAIL is primarily related to canopy architecture and soil/vegetation contrast. This link is key to simultaneous estimation of canopy biophysical/structural variables for applications in agriculture, plant physiology, and ecology at different scales.

### C. Experiment 3: Causation in RTM Emulation

As observed before, noise plays a fundamental role in causal discovery. In this experiment we studied the impact of other types of distortions in remote sensing data. In particular, we aim to assess the non-linearities introduced when approximating a physical model via machine learning. This form of surrogate modelling is known in the literature as *emulation*, and has captured the attention in recent years because it allows to replace RTMs with more efficient statistical algorithms [33], [48]. Emulators, however are just function approximators and the simulated radiances are subject to complicated distortions. We evaluate the identifiability of the causal links in such cases.

Here we trained a neural network using the $n = 1000$ points generated by PROSAIL in the previous experiment to build an emulator. Performance showed less than 5% of normalized RMSE in all bands. Once trained, the emulator was run to

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**TABLE II**: Configuration parameters of the simulated data.

| Parameter                                      | Sampling | Min   | Max   |
|------------------------------------------------|----------|-------|-------|
| RTM model: Prospect 4                          |          |       |       |
| Leaf Structural Parameter                      | Fixed    | 1.50  | 1.50  |
| $C_{ab}$, chlorophyll a+b [µg/cm$^2$]           | $t(14,49)$| 0.067 | 79.97 |
| $C_w$, equivalent water thickness [mg/cm$^2$]   | $t(10,31)$| 2     | 50    |
| $C_m$, dry matter [mg/cm$^2$]                   | $t(5.9,19)$| 1.0   | 3.0   |
| RTM model: 4SAIL                                |          |       |       |
| Diffuse/direct light                           | Fixed    | 10    | 10    |
| Soil Coefficient                               | Fixed    | 0     | 0     |
| Hot spot                                       | Fixed    | 0.01  | 0.01  |
| Observer zenith angle                          | Fixed    | 0     | 0     |
| LAI, Leaf Area Index                           | $t(1.2,4.3)$| 0.01 | 6.99  |
| LAD, Leaf Angle Distribution                   | $t(28,51)$| 20.04 | 69.93 |
| SZA, Solar Zenit Angle                         | $t(8.5,31)$| 0.082 | 49.96 |
| PSI, Azimut Angle                              | $t(30,100)$| 0.099 | 179.83 |

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Footnote: http://teledetection.ipgp.jussieu.fr/prosail/
generate \( n = 10^6 \) samples. The same amount of 182 cause-effect bivariate problems as before was generated. We run the different criteria training the regression models with different amounts of data points, \( n = \{10000, 20000, 30000\} \), and the standard AUC and PR criteria are shown in Fig. 9. Results show that (1) all criteria improve performance with an increasing amount of data, and (2) our criterion \( C_s \) outperforms the state-of-the-art \( C \) in all cases (by around +2.4\%).

**D. Experiment 4: Impact of the regression model**

In this last experiment, we are concerned about the use of different regression algorithms that better account for noise and non-linearities [33]. In particular, we will compare the use of the standard (homoscedastic) GP regression model (GP) [32] (cf. Section II), with the heteroscedastic GP model (VHGP) introduced in [33], [49] (which accounts for signal-to-noise relations), and a warped GP model (WGP) introduced in [50], [51] (which further transforms model’s output to look more like a Gaussian process).

We exemplify these different approaches in a relevant geoscience problem. Terrestrial ecosystems absorb approximately 120 Gt of carbon annually from the atmosphere, about half is returned as plant respiration and the remaining 60 Gt yr\(^{-1}\) represent the Net Primary Production (NPP). Out of this, about 50 Gt yr\(^{-1}\) are returned to the atmosphere as soil/litter respiration or decomposition processes, while about 10 Gt yr\(^{-1}\) results in the Net Ecosystem Production (NEP). The problem here deals with estimating the causal relation between the photosynthetic photon flux density (PPFD), which is a measure of light intensity, and the NEP, which results from the potential of ecosystems to sequester atmospheric carbon. Discovering such relations may be helpful to better understand the carbon fluxes and to establish sinks and sources of carbon. We use here three data sets taken at a flux tower at site DE-Hai involving PPFD(total), PPFD(diffuse), PPFD(direct) drivers and the NEP consequence variable [52].

Results for all three scenarios are shown in Table III. We show the values of HSIC in forward and backward directions, as well as the criteria obtained by all regression models. The heteroscedastic GP accounts for the signal-to-noise relations in a more sensible way, so the dependency estimate becomes slightly more reliable. Nevertheless performance of WGP excels in these particular problems, probably because of the better estimation of conditionals in higher density regions (see Fig. 1 in [53]). Future work will involve testing these models in a wider range of applications.

**V. CONCLUSIONS**

This paper introduced for the first time the issue of observation-based causal inference in bivariate instantaneous problems in remote sensing and geosciences. Approaching this kind of problems requires taking some (strong) assumptions, so results must be taken with extreme caution. Nevertheless, the obtained results confirm that in general causal detection accuracy is well above chance, and opens the field to further experimentation.

To tackle this challenging problem, we used a simple method based on regression and dependence estimation, and proposed a new criterion based on the sensitivity (derivative) of the dependence estimator instead of the dependence itself. This allows us to better capture the asymmetry of the forward and inverse densities with regard to the causal mechanism.

State-of-the-art accuracy was obtained in a wide range of situations with known ground truth. We illustrated performance in a collection of 28 geoscience causal inference problems, in a large database of PROSAIL simulations and emulators in vegetation parameter modeling, and in a carbon

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**TABLE III: Results in the ‘PPFD causes NEP’ causal problem.**

| Method | HSIC\(_f\) | HSIC\(_b\) | C\(_f\) | C\(_b\) | Conclusion |
|--------|-----------|-----------|--------|--------|------------|
| GP     | 6.7525    | 10.3255   | 3.7729 | -1.9294| PPFD(diff)→NEP |
| VHGP   | 6.8220    | 11.7661   | 4.9441 | -1.8758| PPFD(diff)→NEP |
| WGP    | 6.8670    | 12.0412   | 5.1742 | -1.6784| PPFD(diff)→NEP |

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\(^5\)The total PPFD was measured here as the number of photons falling on a one square meter area per second, while NEP was calculated by photosynthetic uptake minus the release by respiration, which is known to be driven by either the total, diffuse or direct PPFD.
cycle problem. We evaluated the impact of using different regression models based on Gaussian Processes; as well as assessed identifiability in the presence of different noise sources and distortions. Models performance was evaluated in global terms by measuring the right direction of causation using standard metrics derived from detection curves.

We would like to finally note that the methodologies proposed here were originally introduced for general-purpose applications. We have nevertheless shown its applicability in remote sensing and geosciences. We relied on a general assumption of structural models in general and ANM in particular. If the assumptions are not fulfilled, the method should not perform well. This may happen in some cases, such as in cases of post-nonlinear effects. Actually, we showed this case experimentally where several regression models were used. The WGP generalizes does not assume an additive noise model in general and results actually confirm that, by replacing the regression model, one can achieve more robust results in cases where assumptions are not met.

The proposed scheme for bivariate causal inference can actually include many independence criteria, such as e.g. differential entropies, empirical Bayes scores or minimum message length scores. We however restricted ourselves to the (standard use of) HSIC, and included a novel causal criterion (the sensitivity map) that can be computed analytically from HSIC. Other more sophisticated criteria could be actually derived from the combination of the sensitivity map and the HSIC values and $p$-values, which will be matter of future research.

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