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A Flexible Smoother Adapted to Censored Data With Outliers and Its Application to SARS-CoV-2 Monitoring in Wastewater

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Courbariaux M, Cluzel N, Wang S, Maréchal V, Moulin L, Wurtzer S, Obépine Consortium, Mouchel J-M, Maday Y and Nuel G (2022) A Flexible Smoother Adapted to Censored Data With Outliers and Its Application to SARS-CoV-2 Monitoring in Wastewater. Front. Appl. Math. Stat. 8:836349. doi: 10.3389/fams.2022.836349 A sentinel network, *Obépine*, has been designed to monitor SARS-CoV-2 viral load in wastewaters arriving at wastewater treatment plants (WWTPs) in France as an indirect macro-epidemiological parameter. The sources of uncertainty in such a monitoring system are numerous, and the concentration measurements it provides are left-censored and contain outliers, which biases the results of usual smoothing methods. Hence, the need for an adapted pre-processing in order to evaluate the real daily amount of viruses arriving at each WWTP. We propose a method based on an auto-regressive model adapted to censored data with outliers. Inference and prediction are produced *via* a discretized smoother which makes it a very flexible tool. This method is both validated on simulations and real data from *Obépine*. The resulting smoothed signal shows a good correlation with other epidemiological indicators and is currently used by *Obépine* to provide an estimate of virus circulation over the watersheds corresponding to about 200 WWTPs.

Keywords: measurement error, smoothing algorithm, outliers, censored data, SARS-CoV-2, autoregressive model

1. INTRODUCTION

A sentinel network, *Obépine (Observatoire épidémiologique dans les eaux usées)*¹, has been designed to monitor SARS-CoV-2 viral load in wastewaters arriving at about 200² wastewater treatment plants (WWTPs) in France as an indirect macro-epidemiological parameter. This survey was initiated in March 2020 in WWTPs of the Greater Paris, during the first epidemic wave in France [1]. In this system, samples are currently typically taken two times a week, resulting in missing data on most days.

The sources of uncertainty in such monitoring system are numerous [2, 3]. These include, for example:

¹https://www.reseau-obepine.fr/

²In November 2021, representing more that one third of the French population.

- sampling variance (although sampling at the WWTPs is integrated over 24 h),
- variance from the Reverse Transcriptase (RT) and quantitative or digital Polymerase Chain Reaction (qPCR or dPCR) processes used to measure the concentration of SARS-CoV-2 RNA genes in the samples,
- uncertainties on other analytical steps in the labs such as virus concentration, genome extraction, and the presence of inhibitors.

Using the raw viral load measurements right away to monitor the pandemic can, therefore, be misleading, as a large variation in the measured concentration can either be due to a real variation in virus concentration or to a quantification error. Therefore, these data need pre-processing in order to provide an accurate estimate of the real daily amount of virus arriving at each WWTP and to evaluate the uncertainty on this estimate as also underlined by [3], who tested, for this purpose, 13 concurrent smoothing methods on data from 4 Austrian WWTPs.

A solution to estimate the underlying true concentrations (from which the noisy measurements are derived) is to exploit the time dependence of the successive measurements. This temporal dependence is indeed due to this (never observed) underlying process, while the measurement noises from one time step to another are assumed to be independent. A natural way of exploiting this time dependence to denoise the signal is Kalman filtering [4, 5]. It is one of the methods that [6] used for concentration monitoring, as well as one of the methods experimented by [3].

Concentration measurements provided by the Obépine network are left-censored because of detection and quantification limits in (RT-)qPCR [a problem pointed out by 7, for instance]. This drawback, which is not taken into account in the study of [3], results in a non-linear dependence between the measurements and the underlying process which cannot be handled by the classic Kalman filter. Such censoring obstacle to Kalman filtering can also happen in many other fields (such as visual tracking, due to camera frame censoring, refer to [8], for instance). The Ensemble Kalman (EKF) permits handling this problem by local linearization, but this linearization is sometimes not accurate enough which can lead to large errors [9]. Unscented Kalman Filter [UKF, 10] offers a solution to this problem with an enhanced statistical linearization around the state estimates. The Particle Filter [PF, 11] also permits handling censored data but with a high computational cost. Finally, the Tobit Kalman filter [12] is designed to handle censoring and has a reduced computational cost compared to PF [13]. In this approach, the censored data are replaced within an Expectation-Maximization algorithm by their conditional expectation with the current parameters. However, we wish not only to predict in real time the value of the underlying process being measured but also to reconstruct its past values and to estimate the error of the measurement system. To do so, a backward filter has to be computed and superimposed on the previously described (forward) filters, resulting in smoothers [14].

Data from the *Obépine* network moreover contains numerous outliers (which may be caused by the vagaries of sampling or by a handling error, for example) that can bias a smoothing that would treat them as regular measurements. [15] review adaptations of Kalman smoothing robust to such outliers (and also to abrupt changes of the underlying state and to switching linear regressions). This is mainly done by modifying the loss function to be minimized. However, those adaptations do not apply to the censoring scheme.

In this study, we here focus on the one-dimensional setting and propose a Smoother adapted to Censored data with OUtliers (SCOU) that answers both the censoring and the outliers detection problems through discretization of the state space of the monitored quantities and more generally permits a very high degree of modeling flexibility.

The proposed model and its implementation are described in section 2. Section 3 provides an illustration and validation of our approach using numerical simulations. Section 4 gives an example of utilization of data from the *Obépine* network.

2. THE PROPOSED METHOD

2.1. An Auto-Regressive Model

Our method is based on the following state-space model:

$$X_t = \eta X_{t-1} + \delta + \sigma \varepsilon_{X,t}$$

$$O_t \sim \mathcal{B}(p)$$

$$(Y_t^* | O_t = 0) = X_t + \tau \varepsilon_{Y,t}$$

$$(Y_t^* | O_t = 1) \sim \mathcal{U}([a, b]) \qquad (1)$$

$$Y_t = \max(Y_t^*, \ell)$$

$$\begin{pmatrix} \varepsilon_{X,t} \\ \varepsilon_{Y,t} \end{pmatrix} \stackrel{i.i.d}{\sim} \mathcal{N}(0, I),$$

where:

- *t* is the time index (ranging from 1 to *n*).
- X_t ∈ ℝ is the real quantity at time t and X = (X_t)_{t∈{1,...,n}} is the vector of real quantities (to be recovered).
- $Y_t \in \mathbb{R}$ is the measurement at time t. Y_t is generally only partially observed. We note $\mathcal{T} \subset \{1, ..., n\}$ the set of t at which Y_t is observed. $Y = (Y_t)_{t \in \mathcal{T}} = Y_{\mathcal{T}}$ is the vector of measurements. Y^* is an accessory latent variable corresponding to a non-censored version of Y. I is the identity matrix.
- $\eta \in \mathbb{R}, \delta \in \mathbb{R}, \sigma \in \mathbb{R}^+, p \in [0, 1] \text{ and } \tau \in \mathbb{R}^+ \text{ are parameters}$ (to be estimated).
- ℓ is the threshold below which censorship applies³.
- $O_t \in \{0, 1\}$ is, for any $t \in \mathcal{T}$, the indicator variable of the event " Y_t^* is an outlier". $O = (O_t)_{t \in \mathcal{T}}$. $\mathcal{B}(p)$ stands for the Bernouilli distribution of parameter p and $\mathcal{U}([a, b])$ for the Uniform distribution on the interval [a, b]. a and b have to be chosen, they can for example correspond to quantiles (respectively, very close to 0 and very close to 1) of the empirical marginal distribution of Y.

 $^{^3}$ In practice, ℓ can vary from one day to another, for instance if one works on quantities that correspond to the multiplication of concentrations (with a detection limit) by a fluctuating volume. This can be taken into account within our method with no additional cost.



Figure 1 depicts this model. In this figure, the arrows represent the direct dependency relations between the variables. We can see the auto-regressive nature of the latent process X, while the Y is independent conditionally to X being known.

2.2. Inference and Prediction—Smoothing and Discretization

We are interested in the distribution of $(X_t|Y_T = y_T)$ for every $t \in \{1, ..., n\}$.

2.2.1. The Smoother

This distribution can be computed from the forward (F) and backward (B) quantities, which are defined as follows:

• *F* is a (Kalman) Filter. For t = 1, ..., n,

$$F_t(x) = \mathbb{P}(X_t = x, Y_{\{1, \dots, t\} \cap \mathcal{T}} = y_{\{1, \dots, t\} \cap \mathcal{T}}).$$
(2)

• $B_n(x) = 1$ (by convention). For t = n - 1, ..., 1,

$$B_t(x) = \mathbb{P}(Y_{\{t+1,\dots,n\} \cap \mathcal{T}} = y_{\{t+1,\dots,n\} \cap \mathcal{T}} | X_t = x).$$
(3)

For $t \in \{1, ..., n\}$, $F_t(x)B_t(x) \propto \mathbb{P}(X_t = x|Y_T = y_T)$. In the strictly Gaussian case, the F and B quantities can be computed recursively in closed formulas. Censorship and outliers, unfortunately, prevent the use of these closed formulas.

2.2.2. Discretization

We have, hence, developed a discrete version of those computations, which makes it easy to adapt to many possible extensions of the model (censorship but also outliers and possibly heteroscedasticity for instance). The set of values that can be taken by *X*, approximated by [*a*, *b*], is discretized into *D* values and becomes \mathcal{X} , with, for example, $\mathcal{X} = \{a, a + \Delta, a + 2\Delta, ..., b\}$ and $\Delta = \frac{b-a}{D-1}$. We then return to the framework of hidden

Markov chains with D states, the D hidden states being the D possible values for each X.

The transition matrix, π , is calculated as follows:

$$\pi(x,x') = \mathbb{P}(X_{t+1} = x' | X_t = x) = \frac{f_{\mathcal{N}}(x'; \eta x + \delta, \sigma^2)}{\sum_{x'' \in \mathcal{X}} f_{\mathcal{N}}(x''; \eta x + \delta, \sigma^2)},$$

where $f_{\mathcal{N}}(.; \mu, \sigma^2)$ is the Gaussian distribution function with mean μ and variance σ^2 .

Let $e_t(x) = \mathbb{P}(Y_t = y_t | X_t = x)$. For all $x \in \mathcal{X}$, $e_t(x) = 1$ if $t \notin \mathcal{T}$. If $t \in \mathcal{T}$:

$$e_{t}(x) = (1-p) \left[\mathbb{I}_{\{y_{t} > \ell\}} f_{\mathcal{N}}(y_{t}; x, \tau^{2}) + \mathbb{I}_{\{y_{t} \le \ell\}} F_{\mathcal{N}}(\ell; x, \tau^{2}) \right] \\ + p \left(\mathbb{I}_{\{y_{t} > \ell\}} \frac{1}{b-a} + \mathbb{I}_{\{y_{t} \le \ell\}} \frac{\ell-a}{b-a} \right),$$

where $\mathbb{I}_{\{A=a\}}$ is the indicator variable for the event $\{A = a\}$ and $F_{\mathcal{N}}(.; \mu, \sigma^2)$ is the Gaussian cumulative distribution function with mean μ and variance σ^2 .

The F and B quantities, defined in Equations (2) and (3), can then be calculated recursively from π and e:

• $F_1(x) = \frac{e_1(x)}{D}$ and, for $t \in \{2, ..., n\}$,

$$F_t(x') = \sum_{x \in \mathcal{X}} F_{t-1}(x)\pi(x, x')e_t(x').$$

• $B_n(x) = 1$ (by convention) and, for $t \in \{n, ..., 2\}$,

$$B_{t-1}(x) = \sum_{x' \in \mathcal{X}} \pi(x, x') e_t(x') B_t(x').$$

Numerical tricks like logarithmic re-scaling are moreover used to calculate e, π and the F and B quantities in order to avoid underflow and overflow problems.

2.2.3. Learning

The parameters η , δ , σ , τ , and p maximize the likelihood, which is given by $\sum_{x \in \mathcal{X}} F_n(x)$, could be obtained with the Expectation-Maximization algorithm. However, it is made challenging by the discretization and renormalization. This is why numerical optimization is preferred. The parameters are more precisely obtained by the numerical optimization of Nelder and Mead (as implemented in the R function optim).

2.2.4. Prediction

For any *t*, the marginal distribution (mean, SD) of $(X_t|Y_{\mathcal{T}} = y_{\mathcal{T}})$, is given by $F_t(x)B_t(x) = \mathbb{P}(X_t = x, Y_{\mathcal{T}} = y_{\mathcal{T}}) \propto \mathbb{P}(X_t = x|Y_{\mathcal{T}} = y_{\mathcal{T}})$ (proof⁴ left to the reader) that is, for any $x \in \mathcal{X}$, the probability that X_t (once discretized) takes the value *x* conditionally to $Y_{\mathcal{T}} = y_{\mathcal{T}}$.

2.2.5. Simulations

We similarly show that:

$$\mathbb{P}(X_{t-1} = x, X_t = x', Y_{\mathcal{T}} = y_{\mathcal{T}}) = F_{t-1}(x)\pi(x, x')e_t(x')B_t(x').$$

⁴Hint: decompose $(Y_{\mathcal{T}} = y_{\mathcal{T}}) = (Y_{\{1,\dots,t\}} \cap \mathcal{T} = y_{\{1,\dots,t\}} \cap \mathcal{T}, Y_{\{t+1,\dots,n\}} \cap \mathcal{T} = y_{\{t+1,\dots,n\}} \cap \mathcal{T})$, condition on $X_t = x$ and use the conditional independence.

Thus, to simulate from $\mathbb{P}(X_{\{1,\dots,n\}}|Y_{\mathcal{T}} = y_{\mathcal{T}})$, one can process sequentially, simulating x_1 with $\mathbb{P}(X_1 = x|Y_{\mathcal{T}} = y_{\mathcal{T}}) \propto F_1(x)B_1(x)$, then, for $t \in \{2,\dots,n\}$:

$$\mathbb{P}(X_t = x' | X_{t-1} = x, Y_{\mathcal{T} = y_{\mathcal{T}}}) = \frac{\pi(x, x')e_t(x')B_t(x')}{B_{t-1}(x)}$$

2.2.6. Outliers Detection

The probability for each observation Y_t to be an outlier (knowing $Y_T = y_T$ and for the *a priori* probability of *p*) can be computed as follows:

$$\mathbb{P}(O_t = 1 | Y_{\mathcal{T}} = y_{\mathcal{T}}) = \frac{p \mathbb{P}(Y_{\mathcal{T}} = y_{\mathcal{T}} | O_t = 1)}{\mathbb{P}(Y_{\mathcal{T}} = y_{\mathcal{T}})}$$
(4)
$$= \frac{p \sum_x \sum_{x'} F_{t-1}(x) \pi(x, x') B_t(x') \left(\mathbb{I}_{\{y_t > \ell\}} \frac{1}{b-a} + \mathbb{I}_{\{y_t \le \ell\}} \frac{\ell-a}{b-a} \right)}{\sum_x \sum_{x'} F_{t-1}(x) \pi(x, x') B_t(x') e_t(x')}.$$

An outlier is detected if $\mathbb{P}(O_t = 1 | Y_T = y_T) > h$, where *h* has to be chosen according to the targeted false positive and false negative rates.

3. NUMERICAL ILLUSTRATIONS ON ARTIFICIAL DATA

The proposed estimation is first evaluated on artificial data. The simulations are designed to study the ability of the algorithm to produce a good estimation of the parameters, to identify correctly the outliers, and to adequately predict the conditional distribution of the underlying process.

3.1. Data Generation and Protocol

Artificial data are simulated according to Model (1) with n = 150 time steps, 50% of which are observed. We set *a* and *b*, respectively, to the quantiles 0.02% and 99.98% of the marginal distribution of Y'. For the sake of consistency with the study presented in section 4, parameters are set so as to be realistic with regard to the *Obépine* data: $\sigma = 0.3$, $\tau = 0.6$, η close to 1 and δ close to 0.

We realize five experiments:

- with no censoring, no outlier, $\delta = 0, \eta = 1$,
 - · (1) with $\Delta = 0.02$,
 - · (2) with $\Delta = 0.1$,
 - $\cdot \quad (3) \text{ with } \Delta = 0.7,$
- (4) with Δ = 0.1, ℓ set for each simulated data such that 16% of Y' are censored (medium censoring level), an outlier rate of p = 7%, η = 0.99 and δ = 0.001,
- (5) same setting as the previous experiment but with 31% of censored data (high censoring level).

Each experiment is replicated 100 times. For each simulation, we compare our approach, SCOU, to:

- a 2-parameter Kalman Smoother implemented in the DLM R package [16]. In the latter method, censoring and outliers are not taken into account and δ and η are not estimated (δ is set to 0 and η is set to 1). Those hypotheses correspond to the settings of the experiments (1), (2), and (3), which aims at checking that the two methods give identical results if Δ is small enough.
- the moving average smoother.
- the Locally Estimated Scatterplot Smoothing (LOESS) [17] which consists of locally weighted polynomial regressions.

As in the study by [3], the parameters of the two last methods (the number of days of the moving average smoother and the span parameter of the LOESS) are chosen by leave-one-out cross-validation so as to minimize the Root Mean Squared Error (RMSE) for the prediction of the left out observations. When (left-)censoring occurs, considering the observed censored values would result in poor performance of those smoothers in recovering the underlying auto-regressive signal. We compensate for this by taking the estimated mean for the corresponding right-truncated normal distribution in ℓ (estimated by assuming that the observations follow a left-censored normal distribution in ℓ) as the observed value for those smoothers, which results in improved performances in all cases.









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3.2. Results on Artificial Data

3.2.1. Impact of Discretization

The first, second, and third experiments (no outliers, no censoring, $\delta = 0$ and $\eta = 1$) show, as expected, that the



FIGURE 5 | Root Mean Squared Errors for the prediction of *X* for 4 competitive smoothing methods [the proposed smoother (SCOU), the 2-parameter Kalman Smoother, the LOESS, and the moving average (MA)], with 16% and 31% of censored data (respectively white and cyan boxes).



2-parameters Kalman smoother implemented in the DLM R package and our method give identical results for $\Delta = 0.02$. The two methods show close performances for $\Delta = 0.1$ and a substantial degradation when $\Delta = 0.7$, as shown in **Figure 2**, where the RMSE (Root Mean Square Error) for the prediction of *X* is computed for each of the 100 repetitions of each of the three experiments.

In the following, we focus on the two other experiments (medium and high censoring levels, p = 7%, $\eta = 0.99$, $\delta = 0.001$, and $\Delta = 0.1$).

3.2.2. Illustration on One Simulation Example

Figure 3 illustrates, on an example, the results of our method on simulated data within the fourth experiment setting (medium censoring level). On this illustration, we can see that the learning process and the use of the smoother permit to finely predict the trajectory of the underlying process, X (in red), from the observations $Y_{\mathcal{T}} = y_{\mathcal{T}}$ (represented by dots) while adequately taking into account the time interval during which the censoring applies. On this figure, the more pink-colored the points are, the higher the estimated probability of them being an outlier. For a detection threshold set lower than h = 0.95, two out of the four simulated outliers (pink crosses and circles, two of them being censored) are identified. The 23 days moving average and the LOESS (for span= 0.24) smoothers, whose parameters are selected by leave-one-out cross-validation, give rather good reconstitutions but fail in reconstructing the trajectory when censoring or outliers happen.

3.2.3. Parameter Estimation

Figure 4 shows that the parameters are correctly estimated with the proposed method. However, one can notice a little negative bias in η estimates which might be due to the asymmetric impact of the estimation bias, with a strong degradation if the η estimate exceeds one. The parameters learned by our method are indeed close on average to the parameters used for the simulation of the data. The parameters learned with the 2-parameter Kalman smoother (σ -dlm and τ -dlm) present an estimation bias which illustrates the necessity to take into account the censoring of the data and the outliers when they exist.

3.2.4. Outlier Identification

The *a posteriori* probability that an observation Y_t is an outlier, $\mathbb{P}(O_t = 1 | Y_T = y_T)$, is computed as specified Equation (4) with, for each simulation, the estimated parameters. The ROC (Receiver Operating Characteristic) curves for the detection of the simulated outliers (around 525 out of about $\frac{n \times 100}{2} = 7500$ observations of Y), across all simulations taken altogether, show correct outlier detection performances despite the censoring of some outliers and the possibility for the outliers to take values very close to Xs (the outliers being simulated from a Uniform distribution). Indeed, the AUC (Area Under the Curve) is 0.74 ± 0.01 for both censoring levels. When the true outlier *a priori* probability, p = 0.07, is provided, they are, respectively, of 0.817 ± 0.014 and 0.767 ± 0.016 for the medium and high censoring settings. Here, the standard deviations for the





computed AUCs are obtained by sampling with discount the 7,500 available observations.

3.2.5. Prediction of the Underlying Process Distribution

Finally, we evaluate the ability of our method to predict the distribution of *X* conditionally on the set of observed *Y*, i.e., the distribution of $(X|Y_T = y_T)$. As illustrated in **Figure 5**, in both the medium and high censoring level experimental settings, the RMSEs obtained by our method are significantly lower than the ones obtained with the 2-parameter Kalman Smoother, with the LOESS method, and with the moving average even when their parameters (respectively, the span and the number of days) are chosen so as to minimize the RMSE obtained by leave-one-out cross-validation.

As for the variance prediction, the coverage rates of the 95% Prediction Intervals of our method, derived from the predicted distributions of $(X_t|Y_T = y_T)$, are (on average) close to the target of 95%, with a median coverage rate of 93% (compared to 91% with the 2-parameter Kalman Smoother) in the medium censoring level setting (respectively, 93% and 85% in the high censoring level setting) as illustrated in **Figure 6**.

4. APPLICATION TO THE DATA FROM OBÉPINE

The developed smoothing method aims to provide an estimate of the actual amount of viral genome arriving at each WWTP and to assess the uncertainty of this estimation.

The concentration measurements provided by *Obépine* are adjusted beforehand for rainfalls and wastewater sources other than from households, which can, otherwise, distort the conclusions when it is abundant by diluting the water arriving at the WWTPs [18].

The direct application of SCOU on the flow-adjusted measurements shows heteroscedasticity of the residuals [estimated by $y_T - \mathbb{E}(X_T | Y_T = y_T)$]: their variance increases with *y* (even after removal of numerous measurements identified as outliers and of censored measurements). Besides, the underlying process, *X*, follows the dynamic of an epidemic. During the exponential growth, it is, thus, supposed to multiply from one day to another. For both those reasons, a logarithmic transformation of the *Obépine* data was performed prior to applying the proposed method.

Hence, for this data set, *Y* is the logarithm of the flow-adjusted measured concentrations, *X* is the logarithm of the actual virus concentrations (to be estimated), ℓ is the logarithm of the limit below which censoring occurs (typically log(1000*U/L*) for the quantification of the SARS-CoV-2 E gene, where U stands for RNA Units, but it can fluctuate from one day to another due to flow adjustments).

The application of SCOU to real data from the *Obépine* network is illustrated for two WWTPs in **Figure 7**. As shown by the successive predictions, once the parameters are fixed, the predictions are rather stable from one day to another.

The estimates of the smoothing parameters for the 190 *Obépine* WWTPs with enough available observations (at least 10 measurements) are illustrated in panel (**A**) of **Figure 8**. The estimated δ parameters are above 0 while η parameters are very close to 1. Hence, when no new information is provided (no new *Y* is observed), the smoother predicts, on average, an increase in *X* values.

The uncertainty of the monitoring system is evaluated by the parameter τ , whose estimates distribution for the 190 *Obépine* WWTPs is illustrated in panel **(B)** of **Figure 8**. The average value for τ is 0.54 log(U/L). This corresponds to an SD of about 0.65*x*, where *x* is the real E-gene RNA concentration (in U/L) to which the SD of the measurement error is supposed to be





wave to the May 21, 2021.

proportional in our model. The allocation of this uncertainty between the sampling error, the RT-qPCR or RT-dPCR error, and other possible sources of error throughout the system is, however, not known.

Importantly, the resulting smoothed signal is well correlated with the logarithm of the local COVID-19 incidence rates, and this correlation is most of the time greatly enhanced by the proposed smoothing step as depicted in **Figure 9** for the 15 cities with enough data available on both sides. On this figure, the correlations are only computed for dates at which raw data was available and correspond to the best correlation for a time lag ranging from 1 to 3 days in one direction or the other. They are computed for a period ranging from the beginning of the second wave to the May 21, 2021. Those correlations are not expected to be higher since, contrary to incidence rates, the indicator also points to asymptomatic people infected by SARS-CoV-2 and is neither biased by people getting tested outside their city (during holidays for instance) nor by varying testing policies.

In order to produce comparable values from one WWTP to another, *Obépine* moreover performs scaling of the data after smoothing [18]. This scaling takes into account, e.g.,

the maximum amount of virus measured during the epidemic wave that occurred in Autumn 2020 in France, the range of volume treated by the WWTP, and the specifics of the laboratory that analyzed the sample by RT-qPCR or RT-dPCR. The final computed indicator (called WWI for WasteWater Indicator) shows good behavior with regard to the corresponding incidence rates [18].

5. DISCUSSION

We developed a method to smooth one dimensional timeseries consisting of successive censored measurements with outliers when the associated measurement uncertainty is not known and the measured quantities have an auto-regressive nature. By discretizing the state space of the monitored quantities, the proposed method has the advantage of being easily adaptable to the specificities of the data (such as measurement censoring and the occurrence of outliers). An experiment on artificial data validates the proposed inference and prediction method. Our method has then been successfully applied to

data generated during the Obépine monitoring of SARS-CoV-2 genome concentration in WWTPs [18]. Importantly, the proposed smoothing procedure enhances the correlation of the data with other epidemiological indicators such as the incidence rate of COVID-19. The time lag between the two signals is moreover just a few days [18], making WWI a credible alternative to the evaluation of the incidence rate through massive individual testing. This approach may be especially relevant if massive testing campaigns become less relevant notably with the advancement of the vaccination campaign and the availability of self-tests to the general public. Both of these factors may indeed induce a progressive but significant decline in participation in testing in a few months and a significant dwindling of the population surveyed to monitor the pandemic, potentially making it even more partial than it is now.

The proposed method could be further developed. First, the underlying *X* process could have a longer time dependency than an AR(1) process. We could, thus, develop an AR(p) version of this method to handle this (with p > 1). Besides, the behavior of the marginal *X* process, and thus its parameters, are expected to change as we move from the propagation of the epidemic stage to a decreasing stage [19]. Joint treatment of the WWTPs timeseries could overcome the lack of individual data to face this problem. One could, for example, automatically detect common breakpoints corresponding to a change in the parameters η or δ . Another possibility would be either to use extrinsic knowledge of the reproduction factor of the epidemic as input data or to add it as a latent variable that slowly evolves from one day to another.

Another way to proceed would be to deduce from other available epidemiological data the shape of the signal to be found in wastewater (and thus an adequate smoothing) based on fine modeling of the whole pathway of SARS-CoV-2 from the human population to wastewater such as the one proposed by [20]. However, such a mechanistic representation includes a large number of unknowns (actual number of infected individuals in the population, rate of RNA degradation in wastewater, etc.) which makes it difficult to exploit for the reconstruction purposes aimed here.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found at: https://www.data.gouv.fr/fr/datasets/surveillance-du-sars-cov-2-dans-les-eaux-usees-1.

THE OBÉPINE CONSORTIUM

The *Obépine* consortium's Scientific Coordination and Steering Committee (CCOS) is composed of Isabelle Bertrand, Mickaël Boni, Christophe Gantzer, Soizick F. Le Guyader, Yvon Maday, Vincent Maréchal, Jean-Marie Mouchel, Laurent Moulin, Rémy Teyssou and Sébastien Wurtzer. The Scientific Interest Grouping (GIS) *Obépine* is directed by Vincent Maréchal associated with two co-directors: Christophe Gantzer and Laurent Moulin. This group includes the CNRS, Eau de Paris, EPHE, Ifremer, Inserm, IRBA, Sorbonne University, Clermont Auvergne University, Lorraine University, and Université de Paris.

AUTHOR CONTRIBUTIONS

YM, J-MM, VM, LM, and SW brought on the scientific problem. MC contributed to the design of the algorithm, performed experiments on artificial and real data, and wrote the manuscript. GN contributed to the design of the algorithm and coordinated the experiments and the writing of the manuscript. NC and SW prepared the data provided by *Obépine*, contributed to the design of the algorithm, performed experiments on artificial and real data, and discussed the results. NC, SW, J-MM, YM, and VM proofread and discussed the content of the manuscript. YM coordinated the exchanges about the article within the *Obépine* consortium. All authors contributed to the article and approved the submitted version.

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