Abstract—We present a method for jointly modeling power generation from a fleet of photovoltaic (PV) systems. We propose a white-box method that finds a function that invertibly maps vector time-series data to independent and identically distributed standard normal variables. The proposed method, based on a novel approach for fitting a smooth, periodic copula transform to data, captures many aspects of the data such as diurnal variation in the distribution of power output, dependencies among different PV systems, and dependencies across time. It consists of interpretable steps and is scalable to many systems. The resulting joint probability model of PV fleet output across systems and time can be used to generate synthetic data, impute missing data, perform anomaly detection, and make forecasts. In this paper, we explain the method and demonstrate these applications.

Index Terms—photovoltaic systems, photovoltaic fleet modeling, distributed power generation, power generation planning, forecasting, convex optimization, copula method, probability distributions, forecast uncertainty

I. INTRODUCTION

Modeling the power output of a fleet of photovoltaic (PV) systems is of great importance for digital operations and maintenance in the PV sector, which is now a multi-billion dollar industry [1]. Applications include predicting power production, detecting anomalies, and making informed decisions about when and where to send workers to service a site. In recent years, there has been a significant increase in the deployment of PV systems, making it necessary to develop scalable models that can handle thousands of systems simultaneously, while staying robust to real-world data challenges, such as the ability to handle missing data.

In this paper we propose a method to estimate the joint probability distribution of the power outputs of a fleet of PV systems, modeling all relevant correlations in the data across individual PV systems and across time. As a copula method, we first develop a novel set of nonlinear marginal transforms that map the power from each system to a scalar Gaussian, and then develop a set of linear transformations that model the marginally transformed data as a large joint Gaussian distribution. We interact with that model to carry out various applications including synthetic data generation, data imputation, anomaly detection, and forecasting.

II. PRIOR WORK

a) Fleet models: Modeling PV systems for operations and maintenance (O&M) purposes based on measured data has a long history [2]–[7]. These techniques focus on predicting either the maximum power point or the full current-voltage relationship of a PV system under a given set of environmental conditions. O&M tasks are then carried out using the model. Fault detection is performed by comparing actual system power generation to the predicted power from the model. Forecasting future PV system power output is done by running the models on predicted weather trends, such as those generated by NOAA [8]. ‘Fleet modeling’ is the practice of constructing an independent, bespoke model for each system in a PV fleet and is quite labor intensive. More recently, researchers have attempted to reduce the effort of fleet modeling using systematic approaches such as learning algorithms and machine inference [9]. These approaches tend to be more task dependent, e.g., focusing on the task of anomaly detection [10]–[12] or forecasting [13]–[19]. While these methods reduce the human effort to create a PV system model, they do not provide joint models of system behavior in a fleet. Several recent papers have explored models to predict aggregate quantities of fleets, such as temporal variability and maximum feed-in power [20]–[23].

b) Copula models and Gaussianization: Our proposed method draws on previous work on both copula models and Gaussianization methods.

Copulas are tools for modeling dependence of several random variables, first proposed by Abe Sklar in 1959 [24] and recently translated into English in [25]. The basic idea is to apply a nonlinear invertible mapping to each component of a random variable so it has some standard distribution such as uniform or Gaussian, and then model the dependence of these transformed variables [26]–[28]. Copulas have been applied in many domains [29]–[33], including PV data analysis [34]. Copula models may be based on theoretical constructions (e.g., the multivariate Gaussian copula) or may be learned directly from data [27]. The quantile transform is a typical choice for data-driven models of the marginal distributions, and various options exist [35], [36]. Our method includes an autoregressive component in the copula model, an approach that has been explored by other authors [37].

An alternative approach to modeling multivariate joint distributions is Gaussianization, also called ‘normalizing flows’. These methods seek an invertible mapping under which the transformed variable has a standard (jointly) Gaussian distribution [38]–[40]. The transformations are typically built in steps, so the transformation is a composition of multiple
transformations, with the distribution of mapped variables getting closer to Gaussian as more layers or steps are added. We can think of a Gaussian copula as a simple first step in such a normalizing flow. While a normalization method can in principle model any probability distribution, Gaussian copula models cannot. On the other hand a Gaussian copula model makes several operations such as conditioning on some known values very easy, involving just basic linear algebra. (These computations can be done for more complex normalizing flows, but they are much more involved, e.g., requiring Monte Carlo or other sampling type methods.)

c) **Modeling via convex optimization:** Our method relies on convex optimization in every step. This guarantees efficient algorithms for finding global solutions [41], and mature tools exist to easily specify convex optimization problems in code [42], [43]. Many traditional statistical models rely on convex optimization for fitting such as regression [44, Chap. 12], auto-regressive (AR) models [44, Ch. 13], and fitting Gaussian distributions to data [41, §3.5], among many others. Our method is inspired by recent work on the trade-off of fit versus roughness in stratified Gaussian models [45], [46], as well as work on convex optimization based signal decomposition [47].

### III. DATA

We will illustrate our method on PV fleet data provided by SunPower Corporation under a nondisclosure agreement. We select six PV systems located in Southern California, with three grouped in Santa Ana, CA and three grouped in the hills to the east in Tustin, CA. The relative locations are shown in figure 1. This choice of system locations was intentional, as we wanted to verify that our model captures similarity of power profiles for nearby systems.

The data consist of 15-minute (average) power values (in kW) for each of the six systems, recorded from 3/1/2017 to 3/31/2017. Figure 2 depicts the power output of the six systems over the three day period between 3/4/2017 and 3/6/2017. At night the power output is zero; during daylight hours, we see different types of power profiles. On 3/6/2017 we see clear-sky behavior, characterized by a smooth increase until noon followed by a gradual decrease until evening. On 3/5/2017, however, we see the power generation curves with multiple dips and peaks throughout the day, which can be attributed to weather factors such as passing clouds. The maximum power output of the systems varies, with system 4 peaking at around 9 kW, and the other systems peaking at around 2 kW.

We denote the data as $y_t \in \mathbb{R}^d$, with $d = 6$, and the time index running from $t = 1$ to $t = T = 2976$. This particular data set does not have any missing data. However, our method gracefully handles missing values, and indeed, relies on this ability to choose hyper-parameters by cross-validation.

We also use data for the following 2 weeks, from 4/1/2017 to 4/14/2017 as our test set for validating our models and applications. This data was not used to fit our model. The test data set has index running from $t = 1$ to $T_{\text{test}} = 1344$.

### IV. METHOD

We propose a method for fitting the given data $y_1, \ldots, y_T$ to a smooth 24-hour-periodic stochastic process. We apply a sequence of three invertible transformations so that the transformed data is approximately a standard Gaussian. These transformations, which respect periodicity and are constructed to vary smoothly across time, are applied in the three steps shown in figure 3. First we use a smooth periodic nonlinear transform to make the data approximately marginally Gaussian. This allows us to model the changing distribution over a day, in addition to the differing maximum values seen across systems. In the second step we use an autoregressive (AR) model to account for dependencies across time. This results in a residual that is approximately uncorrelated across time. Finally, we fit a smooth periodic Gaussian distribution to the residual of the AR model; from this we can whiten the residual so that it is approximately a standard Gaussian. In the language of copula modeling, our first step is our marginal transformation and the final two steps constitute our copula (i.e., ‘linking’) function.

#### A. Fitting a smooth periodic model

Here we describe the general technique, used in steps 1 and 3 of our method, for fitting a smooth $P$-periodic parameter, given by $\theta_1, \ldots, \theta_T \in \Theta \subseteq \mathbb{R}^m$ to some data, where $\Theta$ is a convex set of allowed parameter values. We will use a Fourier series with $K$ harmonics to represent $\theta_t$.

\[
\theta_t = \sum_{k=0}^{K} \left( \cos \left( \frac{2\pi k t}{P} \right) \alpha_k + \sin \left( \frac{2\pi k t}{P} \right) \beta_k \right),
\]

for $t = 1, \ldots, P$, where $\alpha_k, \beta_k \in \mathbb{R}^m$ are the (vector) coefficients that define $\theta_t$.

We take as a measure of smoothness the Dirichlet energy,

\[
\mathcal{D} = \frac{(2\pi)^2}{P} \sum_{k=1}^{K} k^2 \left( ||\alpha_k||_2^2 + ||\beta_k||_2^2 \right).
\]

The loss function has the form

\[
\mathcal{L} = \sum_{t=1}^{T} \ell_t(\theta_t),
\]
where $\ell_t : \Theta \rightarrow \mathbb{R}$ is a convex loss function that depends on some data at time $t$. (If some data are missing, then the sum is only over $t$ for which the data are available.)

Our generic fitting method takes $\theta$ as a solution of the convex optimization problem

$$\begin{align*}
\text{minimize} & \quad \mathcal{L} + \lambda \mathcal{D}, \\
\text{subject to} & \quad \theta_t \in \Theta, \quad t = 1, \ldots, P,
\end{align*}$$

(2)

where $\lambda > 0$ is the smoothing regularizer hyper-parameter. The variables are the $m$-vectors $\alpha_0, \ldots, \alpha_K$ and $\beta_1, \ldots, \beta_K$. This is a convex optimization problem, and readily solved.

This generic fitting method contains the hyper-parameter $\lambda$ (and possibly others), but good values of these can be found automatically using cross-validation [48, §7.10], [44, §13.2], so the method is essentially hyper-parameter free and automatic.

Two steps of our method solve a different instance of the problem (2) with $\lambda$ chosen using cross-validation. This method of fitting a smooth periodic parameter is a special case of a Laplacian regularized stratified model [45], [46] with the underlying graph a cycle representing the periodicity. It can also be thought of as a signal decomposition problem [47].

### B. Marginal transforms

In this first step, we seek continuous increasing functions $\varphi_{t,i} : \mathbb{R} \rightarrow \mathbb{R}$. We allow these functions to change with $t$ but enforce that they are periodic and smooth. Our goal is for the transformed values,

$$x_{t,i} = \varphi_{t,i}(y_{t,i}), \quad \text{for } t = 1, \ldots, T,$$

to have an approximately Gaussian (marginal) distribution for each $i = 1, \ldots, d$. Typical quantile transforms are static; but our method defines a transformation that changes smoothly in time and is periodic.

We carry out this step for each component of the original data, so the method described in this section is carried out separately for each $i = 1, \ldots, d$. To lighten the notation in this section, we drop the component index $i$, and consider the original data $y_t$ to be scalar.

Our first step is to estimate a set of quantiles $0 \leq \eta_1 < \cdots < \eta_r \leq 1$ of the data. By default we take these to be 2nd percentile, the 98th percentile, and the 10 deciles, so $r = 11$

$$\eta = (0.02, 0.10, 0.20, \ldots, 0.80, 0.90, 0.98),$$

but our method is general and any other choice of quantiles could be used. We denote the estimated $\eta_i$ quantile of $y_t$ as $q_{t,i}$, $t = 1, \ldots, T$, $i = 1, \ldots, r$. We assume these are $P$-periodic and smooth.

To estimate these quantiles from the data we use standard quantile regression [49], [50], which relies on the so-called pinball loss, defined as

$$\ell_{\text{pin}}(u; \eta) = \max\{(1 - \tau)u, \tau u\} = (\tau - 1/2)|u| + (1/2)u,$$

for quantile $\tau \in [0,1]$. We take our loss function to be

$$\ell(q_t) = \ell_{\text{pin}}(q_{t,1} - y_t; \eta_1) + \cdots + \ell_{\text{pin}}(q_{t,r} - y_t; \eta_r),$$

(3)
the sum of the pinball losses associated with each of our \( r \) quantiles. To estimate the quantiles we solve the generic problem (2), with loss function (3), and constraint set
\[
\Theta = \{ q \mid q_1 \leq \cdots \leq q_r \},
\]
which enforces that the quantiles are consistent. This simultaneously estimates the \( r \) quantiles of \( y_t \) for each \( t \), with the quantiles being smooth and periodic, and always satisfying the consistency constraint. The hyper-parameter \( \lambda \), which controls how smooth the quantile estimates are, can be chosen automatically via cross-validation.

Given these periodic smooth quantile estimates, we construct nonlinear mappings \( \varphi_t : \mathbf{R} \rightarrow \mathbf{R} \) as continuous piece-wise linear functions, with knot-points given by the quantiles, and values at those points given by the associated value of a standard scalar Gaussian for the same quantiles, i.e.,
\[
\varphi_t(q_{j,i}) = \Phi^{-1}(\eta_{j,i}), \quad j = 1, \ldots, r,
\]
where \( \Phi \) is the cumulative distribution function (CDF) of a standard Gaussian. This gives the marginally transformed data \( x_{t,i} = \varphi_t(y_{t,i}) \).

### C. Autoregressive model

The time series \( x_1, \ldots, x_T \) has entries with approximately standard Gaussian marginal distribution, but there are dependencies between the components, as well as across time. Our next step is to handle the dependency across time. We fit a vector autoregressive (AR) model to the marginally transformed data \( x_t \). The model is
\[
x_t = A_1 x_{t-1} + \cdots + A_M x_{t-M} + v_t,
\]
where \( v_t \) is a process noise or residual, \( M \) is the memory of the AR model, and \( A_1, \ldots, A_M \in \mathbf{R}^{d \times d} \) are the coefficients. We could fit these AR coefficients as smooth and periodic, but we have found that a constant AR model does just as well as the AR model, and
\[
\mathbf{A} \mathbf{v}_t = \mathbf{y}_t,
\]
where \( \mathbf{v}_t \) and \( \mathbf{y}_t \) are periodic, and values at those points given by the associated value of a standard scalar Gaussian for the same quantiles, i.e.,
\[
\varphi_t(q_{j,i}) = \Phi^{-1}(\eta_{j,i}), \quad j = 1, \ldots, r,
\]
where \( \Phi \) is the cumulative distribution function (CDF) of a standard Gaussian. This gives the marginally transformed data \( x_{t,i} = \varphi_t(y_{t,i}) \).

### D. Smooth periodic residual fit

Our last step is fit a smoothly varying periodic Gaussian distribution to the residual \( v_t, v_t \sim \mathcal{N}(\mu_t, \Sigma_t) \), where we assume that \( v_s \) and \( v_t \) are independent for \( s \neq t \). We model \( v_t \in \mathbf{R}^d \) as smooth periodic Gaussian,
\[
v_t \sim \mathcal{N}(\mu_t, \Sigma_t), \quad t = 1, \ldots, T
\]
where \( \Sigma_t \) and \( \mu_t \) are smooth and periodic. We expect \( \mu_t \) to be small.

Our loss \( \ell_t \) will be the negative log-likelihood of the Gaussian model (6), which is
\[
\ell_t = \frac{d}{2} \log(2\pi) - \sum_{j=1}^{d} \log \left( \text{diag} \left( L_t \right) \right) + \frac{1}{2} \| L_t^T v_t - \nu_t \|_2^2,
\]
with variables \( L_t \in \mathbf{R}^{d \times d} \) and \( \nu_t \in \mathbf{R}^d \). Here, \( L_t \) is the Cholesky factor of \( \Sigma_t^{-1} \) and \( \nu_t = L_t^{-T} \mu_t \). This change of variables makes the loss a convex function. We can recover \( \Sigma_t \) and \( \mu_t \) as
\[
\Sigma_t = (L_t L_t^T)^{-1}, \quad \mu_t = L_t^{-1} \nu_t,
\]
and then solve the optimization problem to find \( L_t \) and \( \nu_t \). Here too the smoothness hyper-parameter \( \lambda \) is found by cross-validation.

Our final whitened signal is given by
\[
z_t = \Sigma_t^{-1/2} (x_t - \mu_t) = L_t^T (x_t - \nu_t),
\]
defined when \( x_t \) is. According to our model, these are independent identically distributed (IID) with \( z_t \sim \mathcal{N}(0, I) \).

### E. The whole model

From our first two steps, we see that our model of \( y_t \) is a stationary periodic Gaussian process \( x_t \), mapped entrywise through a smoothly periodic transformation. Using all three steps, we interpret it as IID Gaussians \( z_t \), passed through an AR filter to obtain \( y_t \), and then mapped entrywise.

Such a model allows us to carry out several operations. We can generate samples from the model. We can evaluate the density at a sequence \( y_t \). We can condition on a set of known values of some of the components, as well as computing conditional marginal quantiles for each unknown entry. These allow us to carry out imputation, i.e., guessing missing values, by evaluating the conditional median of a missing entry given the known ones. (We also can get error bars, e.g., the 10th and 90th conditional marginal quantiles.) We can also do anomaly detection, where we detect known entries that do not fit the model. To do this we compute the conditional quantile of each known entry, given the other known entries but not that particular value; conditional quantile values that are either near zero or one are then flagged as suspicious.

These operations (and others) can be carried out for many types of statistical models, for example using Monte Carlo sampling. But due to the specific structure of our model, we can carry them out using simple linear algebra, which makes the operations fast and reliable. Details of how we implement these operations will be given in a forthcoming paper.
V. RESULTS

Here we show the results of our modeling method on the PV data described above, using default parameters. Estimated quantiles for each system are shown in figure 4. Note that the 98th percentile serves as an effective statistical clear-sky model. The estimated quantiles collapse to zero at night as expected. The spread between the upper quantiles is narrower than that of the lower quantiles, especially around noon. Quantiles for each system exhibit distinct characteristics, with the systems physically near each other showing similar shaped quantiles.

A few samples of the associated piecewise linear copula transforms are shown in figure 5. We see that the same power value is mapped to different points based on the time of day and the system. This shows how our time-aware copula transform adapts to both the time of day and the unique characteristics of different systems as opposed to a standard fixed copula transform, which does not.

We use AR memory $M = 3$, with coefficients. We observe several interesting phenomena in these coefficient matrices. First, the entries of $A_1$ are generally bigger than those of $A_2$ and $A_3$, showing that the previous period plays a larger role in predicting the current values than the previous two values. We also see that the diagonal entries are generally larger than the off-diagonal ones, meaning that the previous values for each system play a larger role in predicting the current value than the previous values of the other systems. However, the many non-zero off-diagonal elements in the coefficient matrices show that the predictions for each system do depend on the previous values of the other systems.

Finally we fit a smoothly varying periodic Gaussian distribution to the residuals of the AR model, shown in figure 6. The top plot shows the means, which are indeed small, as expected. The middle plot shows the standard deviations of the residuals. These are smaller than one, which is approximately the standard deviation of the entries of $x_t$, which means we are able to predict the current values using previous values better than simply guessing $x_t = 0$, i.e., treating them as uncorrelated across time. We can see that the residual standard deviations vary considerably across systems and time of day. Roughly speaking, the residual of system 1 has almost twice the standard deviation of the residual of system 6. We also see that the residual standard deviation is smaller at noon than in the morning and afternoon. The bottom plot shows the correlations of selected pairs of residuals. These correlations are generally around 30%, but we can see that systems that are physically near each other are more highly correlated. We can also see variation of the correlation over the day.

VI. APPLICATIONS

A. Generating simulated data

We generate simulated data from our model by simulating data from the periodic Gaussian stochastic process, and then applying the inverse nonlinearities $\phi^{-1}_t$ to the entries of these samples. Figure 7 shows two simulations of fake data for
system 1, with the actual data for the specific day 4/7/2017 shown at top for reference. They appear quite similar.

B. Conditional marginal quantiles

We can compute the marginal quantiles of any entry, conditioned on all other known entries, in time and across systems. When the entry is unknown, this gives us a sophisticated method for imputing or guessing what the missing value might have been. We can use the conditional marginal median (50th percentile) as the imputed value, with the 10th and 90th percentiles defining an uncertainty interval. Figure 8 shows the marginal conditional quantiles for system 1 at 5 times on two days, one clear and one partially cloudy. We observe that the model correctly adapts the uncertainty bounds to the weather, with tighter bounds on clear days. Additionally, we note that the uncertainty bounds are asymmetric, with decreases in output (say, due to clouds) more likely than increases. The predictions themselves, shown as the circle representing the conditional median, are good.

C. Anomaly detection

We can use marginal conditional quantiles to identify anomalous entries in our data. To estimate whether a given known entry is an anomaly we pretend that it is unknown, compute its conditional marginal CDF given all other known values, and evaluate it at the known value. We can flag an entry as anomalous if this quantile value is less than \( \epsilon \) or more than \( 1 - \epsilon \), where \( \epsilon \) is a threshold value such as \( 10^{-2} \). With this threshold, we would expect a false positive rate around \( 2\epsilon \).

To illustrate this, we consider system 2 on 4/1/2017. We introduce synthetic anomalies by perturbing the power values at 8:30, 10:00, 11:30, 13:00, 14:15 and 15:30, by randomly increasing or decreasing the true values by 15%. Table I shows true values, perturbed values and conditional marginal quantiles of perturbed values, clipped to the range \([0.0001, 0.9999]\).

With threshold \( \epsilon = 0.01 \), we detect all of the artificial anomalies. We also have three false positives, i.e., times when a true value is flagged as an anomaly. The conditional marginal quantiles for this day are shown in figure 9. The vertical axis shows \( \min\{q, 1 - q\} \), with the threshold \( \epsilon = 0.01 \) shown as the darker horizontal line. True negatives are shown as blue circles, and true positives are shown as blue squares. False positives are shown as orange circles. The three false positives are all cases where the true power was low compared to our predicted median. This is not surprising; clouds can easily reduce power output unexpectedly by 15% or more.

D. Forecast

Here we forecast the values of system 2 from 13:15 on, using data from all systems up through 13:00. We show the forecast, which is the median or 50th conditional marginal quantile, along with the conditional marginal 10th and 90th quantiles, which give us confidence bands for the forecast values. This is illustrated in figure 10 on the clear day 4/9/2017 and the cloudy day 4/6/2017. Our forecast on the clear day is very good, with tight uncertainty bands. Our forecast on the
cloudy day is reasonable, but with much wider uncertainty bands.

In addition to forecasting marginal quantiles of a single system, we can generate joint forecasts using all systems. We illustrate this in figure 11, where we forecast the values of system 2 from 13:15 on, using data from all systems up through 13:00. We show three different forecasts, sampled from the full joint conditional distribution. We see that the forecasts agree with the marginal forecasts, in the sense that the forecasts and the cloudy day 4/6/2017. We forecast from 13:15 on, given values for all systems up through 13:00.

VII. CONCLUSIONS

We presented a novel approach to modeling and analyzing fleets of PV systems based on a smooth periodic Gaussian copula transform, and illustrated some of its applications. While we have demonstrated the method on a small example, it can scale gracefully to much larger problems; details will be given in a forthcoming paper.

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