

Regionalization in a global hydrologic deep learning model: from physical descriptors to random vectors

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Abstract

Streamflow prediction is a long-standing hydrologic problem. Development of models for streamflow prediction often requires incorporation of catchment physical descriptors to characterize the associated complex hydrological processes. Across different scales of catchments, these physical descriptors also allow models to extrapolate hydrologic information from one catchment to others, a process referred to as “regionalization”. Recently, in gauged basin scenarios, deep learning models have been shown to achieve state of the art regionalization performance by building a global hydrologic model. These models predict streamflow given catchment physical descriptors and weather forcing data. However, these physical descriptors are by their nature uncertain, sometimes incomplete, or even unavailable in certain cases, which limits the applicability of this approach. In this paper, we show that by assigning a vector of random values as a surrogate for catchment physical descriptors, we can achieve robust regionalization performance under a gauged prediction scenario. Our results show that the deep learning model using our proposed random vector approach achieves a predictive performance comparable to that of the model using actual physical descriptors. The random vector approach yields robust performance under different data sparsity scenarios and deep learning model selections. Furthermore, based on the use of random vectors, high-dimensional characterization improves regionalization performance in gauged basin scenario when physical descriptors are uncertain, or insufficient.

1 Introduction

In hydrology, streamflow prediction is essential for the forecast of water supply, floods, and droughts. It is a challenging task because of interacting hydrological processes (Beven, 1989, 1987; Freeze & Harlan, 1969; Freeze, 1974), spatial-varying parameter uncertainties (Beven & Binley, 1992), and limited observations (Blöschl & Sivapalan, 1995). These challenges have motivated the advancement of hydrologic models from simple to complex. Encompassing more underlying hydrological processes, a complex hydrologic model includes more hydrologic parameters and detailed catchment physical descriptors to address the complexities (Beven, 2001, 2002) and associated scaling issues (McDonnell et al., 2007). But parameterizing such a complex hydrologic model for any individual catchment becomes difficult when hydrologic data are unavailable. Thus, regionalization, which is defined as “how to extrapolate hydrologic information from one area to another” (Blöschl & Sivapalan, 1995), specifies a research topic of modeling catchment runoff prediction using hydrologic information from multiple catchments, which will be given a brief background review in section 2.

Regionalization heavily relies on physical descriptors, such as, soil porosity, catchment elevation, etc. These physical descriptors account for hydrologic complexities and regional differences and are thus intensively used in regionalized hydrologic models, either process-based or data-driven.

Recently, Kratzert et al. (2019a) have presented a regionalized data-driven hydrologic model that greatly outperforms local process models. Specifically, they trained a single deep learning model (LSTM, abbreviated for the Long Short-Term Memory networks) for 531 basins in the US CAMELS (Catchment Attributes and Meteorology for Large Sample studies) dataset (Addor et al., 2017) and show that it is able to greatly outperform the well-established process-based models (e.g., SAC-SMA (Burnash, 1995), VIC (Liang et al., 1994), etc) that have been individually parameterized for each basin, and thus offer a better route to regionalization (Kratzert et al., 2019a).

Building such a model requires streamflow observation and weather forcings for many basins with diverse physical descriptors. It also relies upon the fact that all relevant basin physical descriptors are available and of high quality. Performance of such models may suffer if some of the descriptors are missing or are incorrect/uncertain. Our paper presents an approach where it is possible to build a data driven regionalized model even in the absence of any basin specific physical descriptors. It is able to use the weather forcing and streamflow data from a set of basins to build a global model without having any information about the physical descriptors of individual basins (For the background information of the global model, please see section 2). However the structure of this model is identical to the one used by Kratzert et al.(2019a), as it only replaces the individual catchment physical descriptors by random vectors that simply provide a distinct characterization to each basin. Our results show that this approach provides global models at least as good as the ones produced using the knowledge of all available physical descriptors. But the performance is much better relative to the scenario where some of physical descriptors are missing and/or are incorrect/uncertain.

We note that the random vector and physical descriptor approaches are not in conflict and in fact give comparable results. In fact, for ungauged basins, Kratzert et al.’s model can be used (Kratzert et al., 2019b) and shows that physical descriptors serve as a bridge between gauged basins and ungauged basins. In our approach, the random vectors do not connect gauged basins and ungauged basins due to the lack of streamflow observation for the ungauged basins.

The paper is organized as follows. Section 2 introduces relevant background information, in particular the regionalization. Section 3 explains the details of the random vector method as well as the deep learning architecture involved. This section also explains the dataset and the set up of the experiment. The experiment includes an exhaustive analysis on the applicability of our proposed random vector methods under various data scarce situations and modeling structures. Section 4 lists our benchmarking results and the exhaustive analysis of the random vector applicability. Section 5 highlights scientific implications from our results and suggests a few future directions. Section 6 summarizes the scientific conclusions.

2 Background

Performing hydrologic prediction from multiple catchments, regionalization is closely related to the problem addressed in “prediction in ungauged basins” (PUB) (Sivapalan et al., 2003), and most literature uses “PUB” and “regionalization” interchangeably (Pagliero et al., 2019; de Lavenne et al., 2019; Choubin et al., 2019; Ecrepont et al., 2019; Zamoum & Souag-Gamane, 2019; Prieto et al., 2019; Guo et al., 2021; Alipour & Kibler, 2018). An underlying assumption behind regionalization is that similar basins have similar hydrologic behaviors. This implies that differences/similarities across catchments can be

classified into physical descriptors such as, climatology, geology, geomorphology, etc, with the assumption that incorporating these descriptors will improve streamflow prediction. In other words, hydrological behaviors as predicted from models for different catchments shall be based on similarities with regional information that is characterized by catchment physical descriptors. These approaches have been given a comprehensive review in particular for PUB (Guo et al., 2021; Samaniego et al., 2017; Beck et al., 2016) and can be grouped into model-dependent (process-driven) and model-independent (data-driven) methods, where 'model' denotes process-based models (Prieto et al., 2019).

Model-dependent methods give hydrologic predictions from process-based models. Information from the existing process-based hydrologic model is transferred to ungauged catchments based on certain criteria that link gauged to ungauged catchments. In practice, since those existing hydrologic models are calibrated to a specific catchment, this relies on some strategy of information transfer. A typical application of a model-dependent method implements a well-calibrated local hydrological process-based model and appropriate connections among catchments. In the review paper by Guo et al. (2021), model-dependent methods can be classified into three categories: similarity based methods, regression based methods, and hydrological signature-based methods or some hybrid of each.

The model-independent approaches are data driven and do not rely on physical processes to simulate streamflow. Data driven methods learn how to predict streamflow from weather drivers and catchment physical descriptors directly without involving any hydrological process descriptions. Depending on either one or multiple catchments of data used, the data driven model will learn localized or regionalized hydrologic behaviors respectively. A local model is referred to as the model using hydrologic data from only one catchment. By contrast, when the hydrology data from multiple catchments are used and those catchments cover a wide range of all available hydrologic behaviors, the model is called a global model.

For data driven methods, one family is the neural network (Besaw et al., 2010; Hsu et al., 1995). Besaw built an artificial neural network on one catchment and transferred to another similar catchment without adaptation. It yielded unsatisfactory predictive performance (Besaw et al., 2010). In recent years, the Long Short-Term Memory (LSTM) networks (Hochreiter & Schmidhuber, 1997), one sub-family of neural networks, have shown burgeoning applicability in streamflow prediction tasks (Kratzert et al., 2018). LSTM based methods predict streamflow from antecedent weather drivers. Kratzert et al. (2019a) have shown that using physical descriptors will train a universal global LSTM based model that outperforms process-based individual models given the same forcing data. One of the two versions of the LSTM developed by Kratzert et al. provides additional physical interpretation, that is, basin similarities are preserved in the well trained machine learning (ML) model. In gauged scenarios, Feng (Feng et al., 2020) embedded a global LSTM within a data integration framework (using predicted discharge from previous day) and found that it could marginally reduce prediction bias in regions with high flow autocorrelation. Frame showed that global LSTM outperforms the National Water Model (NWM) (Frame et al., 2020). In the poorly gauged scenarios, Ma (Ma et al., 2021) showed that fine tuning a global LSTM learned from data rich basins improved predictive performance in poorly gauged basins in contrast to local models learned solely from limited data.

It bears emphasis that regionalization approaches, either model-dependent and model-independent, rely heavily on physical descriptors. However, to obtain a satisfactory regionalization performance, physical descriptors need to be sufficient such that process complexities and associated scaling issues (Blöschl & Sivapalan, 1995) are encompassed. Otherwise, catchment scale prediction will be handicapped by the lack of sufficient information. For instance, modeling hydrological behaviors at the small scale can be accomplished by incorporating local processes with a few parameters. However, the incorporated processes and parameterization need to be adjusted, either made simpler or more complex, to model hydrologic behaviors at a larger scale. The same adjustment also oc-

curs when modeling hydrological behaviors between global scale and local scale, upstream and downstream. Accounting for these complexities and heterogeneities, sufficient physical descriptors must be involved. For example, Drost and Mudersbach found that merely incorporating landuse data with no additional physical descriptors provided little improvement to streamflow prediction and therefore may not benefit regionalization (Drost & Mudersbach, 2021). However, due to the complexity of each catchment, such a complete characterization to resolve hydrologic complexity is difficult and challenging (Beven, 2020). This issue will be even more pronounced in applying models to data sparse regions where physical descriptors are limited, or even unavailable.

3 Methods

3.1 Long Short-Term Memory Network

Long short-term memory network (LSTM) (Hochreiter & Schmidhuber, 1997) is a special type of recurrent neural network designed especially for modeling time series predictions. Indeed, LSTM is the state-of-the-art deep learning model to predict streamflow (Kratzert et al., 2018, 2019a; Frame et al., 2020; Feng et al., 2020; Ma et al., 2021). In contrast to a traditional recurrent neural network, LSTM avoids gradient vanishing or explosion (Bengio et al., 1994) and therefore preserves long term temporal dependencies for time series forecasting. This is achieved by using the gating architecture, which explicitly controls information flow and updates system hidden features. This memorizing mechanism and long term dependency allows LSTM to be well suited to model streamflow on a catchment scale. In particular, weather inputs feed and alter catchment response in various temporal scales. Although flooding season yields quick surface water response, the streamflow in winter periods in northern climates tends to have much longer response time because of involved snow and snowmelt processes. With the capability of the LSTM to account for long term dependency, it automatically learns these streamflow behaviors from data. Furthermore, it has been shown that some of the hidden features learned by the LSTM resemble snow processes (Kratzert et al., 2018).

An LSTM maps a sequence of time series input into the response variable. In this paper, we consider an LSTM based architecture that uses input features (\mathbf{x}) spanning T days to predict the observed discharge on the last day of the T -day window. The involved equations of an LSTM models are given below.

$$\mathbf{i}[t] = \sigma(\mathbf{W}_i \mathbf{x}[t] + \mathbf{U}_i \mathbf{h}[t-1] + \mathbf{b}_i) \quad (1)$$

$$\mathbf{f}[t] = \sigma(\mathbf{W}_f \mathbf{x}[t] + \mathbf{U}_f \mathbf{h}[t-1] + \mathbf{b}_f) \quad (2)$$

$$\mathbf{g}[t] = \tanh(\mathbf{W}_g \mathbf{x}[t] + \mathbf{U}_g \mathbf{h}[t-1] + \mathbf{b}_g) \quad (3)$$

$$\mathbf{o}[t] = \sigma(\mathbf{W}_o \mathbf{x}[t] + \mathbf{U}_o \mathbf{h}[t-1] + \mathbf{b}_o) \quad (4)$$

$$\mathbf{c}[t] = \mathbf{f}[t] \odot \mathbf{c}[t-1] + \mathbf{i}[t] \odot \mathbf{g}[t] \quad (5)$$

$$\mathbf{h}[t] = \mathbf{o}[t] \odot \tanh(\mathbf{c}[t]) \quad (6)$$

where $\sigma(\cdot)$ is sigmoid function, $\tanh(\cdot)$ is the hyperbolic tangent function, and \odot means element wise multiplication. \mathbf{W} , \mathbf{U} , \mathbf{b} are model parameters, which will be learned during optimization. Other variables in equations represent basic computation units involved in the calculation. As gating variables, $\mathbf{i}[t]$, $\mathbf{f}[t]$, and $\mathbf{o}[t]$ are input gate, forget gate, and output gate, respectively. They filter the information from the current and the previous time stamp, then combine them to update cell state $\mathbf{c}[t]$. $\mathbf{c}[t]$ underlines the intuition that motivates the LSTM design. $\mathbf{c}[t]$ is maintained serially and embeds the temporal contextual information, which is characterized in $\mathbf{g}[t]$, to then update the hidden representation $\mathbf{h}[t]$. The stacked input \mathbf{x} enters the LSTM sequentially and alters the information inherited from the previous timestamp. The previous information is stored in cell states $\mathbf{c}[t]$ and hidden states $\mathbf{h}[t]$, both of which characterizes the system memory. Cell states $\mathbf{c}[t]$ and hidden states $\mathbf{h}[t]$ are initialized as zero vectors and then grad-

usually modified until the final date in T -day time windows is reached. After a linear transformation, $\mathbf{x}[t]$ infuses with previous hidden state $\mathbf{h}[t-1]$ and then is non-linearly transformed in $\mathbf{i}[t]$, $\mathbf{f}[t]$, $\mathbf{g}[t]$, and $\mathbf{o}[t]$ via a corresponding activation function. The previous timestamp's cell state $\mathbf{c}[t-1]$ is updated with $\mathbf{f}[t]$ and then merges with an element-wise product of $\mathbf{i}[t]$ and $\mathbf{g}[t]$, which injects new information, to form a new cell state $\mathbf{c}[t]$. After another hyperbolic tangent activation, this new cell state $\mathbf{c}[t]$ merges with $\mathbf{o}[t]$ and therefore updates the current hidden state $\mathbf{h}[t]$. After the consecutive alteration of T time stamps, the final hidden state $\mathbf{h}[T]$ is then transformed into the target variable, which in our case is streamflow.

In the context of regionalization based streamflow prediction, both dynamic weather variables and static catchment physical descriptors as formulated in equation 7:

$$Q_t = f(\mathbf{x}^d, \mathbf{x}^s) \quad (7)$$

where Q_t is streamflow, \mathbf{x}^d is weather input vector, and \mathbf{x}^s is a d-dimensional vector of physical descriptors. It bears emphasis that for a given catchment, \mathbf{x}^s is assumed to be temporally static, while \mathbf{x}^d is temporally dynamic. We assume catchment physical descriptors do not vary with the time. In this paper, we consider two widely used LSTM based models as illustrated in Figure 1. Namely, these two models are EA-LSTM and CT-LSTM (Kratzert et al., 2019a), where ‘EA’ denotes entity awareness while ‘CT’ denotes concatenation. These models differ in terms of how \mathbf{x}^s is added into the network. In CT-LSTM physical descriptors are added before LSTM cell, whereas in EA-LSTM, they are used within the cell. For clarifications, the CT-LSTM refers to the normal LSTM used in Kratzert et al.’s paper (2019a). We add prefix ‘CT’ to ‘LSTM’ to emphasize that \mathbf{x}^s is concatenated with weather drivers before entering the LSTM cell.

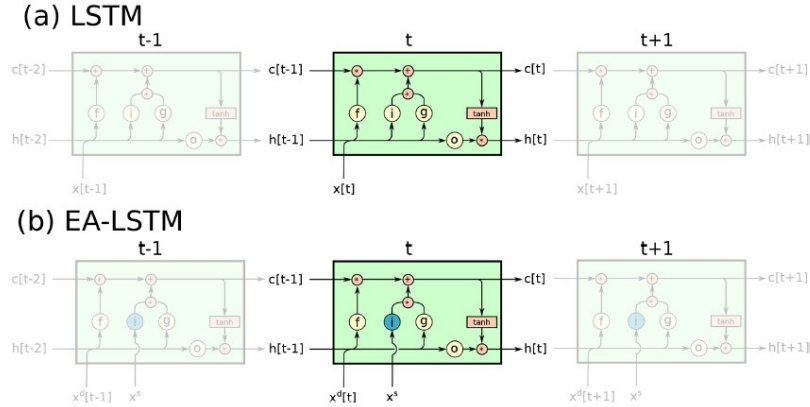


Figure 1: LSTM family illustration. Figure is from “Towards learning universal, regional, and local hydrological behaviors via machine learning applied to large-sample datasets” by Kratzert et al. (2019a), *Hydrology and Earth System Sciences*, 23, 5092 (Kratzert et al., 2019a)

3.1.1 CT-LSTM

In CT-LSTM, at each timestamp, the dynamic weather input \mathbf{x}^d is concatenated with the physical descriptors \mathbf{x}^s to form the model input $\mathbf{x}[t]$:

$$\mathbf{x}[t] = [\mathbf{x}^s, \mathbf{x}^d[t]] \quad (8)$$

This model input enters the LSTM (equation 1 to 6), gets updated via the calculation of gates, and yields the final output - streamflow prediction. Through the calculation, physical descriptors are not placed within the LSTM cells or gates.

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3.1.2 EA-LSTM

First proposed in (Kratzert et al., 2019a), EA-LSTM (Entity Aware LSTM) uses a modified version of LSTM where input gate takes physical descriptors as input instead of input features as previously shown in Equation 1. The key idea here is to explicitly empower the LSTM to customize its learning ability for catchment-wise adaptation.

$$\mathbf{i} = \sigma(\mathbf{W}_i \mathbf{x}^s + \mathbf{b}_i) \quad (9)$$

$$\mathbf{f}[t] = \sigma(\mathbf{W}_f \mathbf{x}^d[t] + \mathbf{U}_f \mathbf{h}[t-1] + \mathbf{b}_f) \quad (10)$$

$$\mathbf{g}[t] = \tanh(\mathbf{W}_g \mathbf{x}^d[t] + \mathbf{U}_g \mathbf{h}[t-1] + \mathbf{b}_g) \quad (11)$$

$$\mathbf{o}[t] = \sigma(\mathbf{W}_o \mathbf{x}^d[t] + \mathbf{U}_o \mathbf{h}[t-1] + \mathbf{b}_o) \quad (12)$$

$$\mathbf{c}[t] = \mathbf{f}[t] \odot \mathbf{c}[t-1] + \mathbf{i}[t] \odot \mathbf{g}[t] \quad (13)$$

$$\mathbf{h}[t] = \mathbf{o}[t] \odot \tanh(\mathbf{c}[t]) \quad (14)$$

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As illustrated in Figure 1b and also equations 9 to 14, \mathbf{x}^s enters the LSTM via input gates, learns customized embedding (equation 9) for each basin, and updates the cell states recurrently at each timestamp. It therefore explicitly controls what modules in LSTM respond to different catchments. This learned embedding will merge with other gates ($\mathbf{f}[t]$, $\mathbf{g}[t]$, $\mathbf{o}[t]$), whose alteration are contributed by only dynamic weather inputs \mathbf{x}^d . This separated role of \mathbf{x}^s and \mathbf{x}^d in EA-LSTM splits the contributions towards stream-flow prediction from \mathbf{x}^s in contrast to \mathbf{x}^d . Additionally, the learned embedding affords an opportunity to examine cross-catchment response in a global model, which was shown to be close to the cross-catchment analysis using true basin characteristics (Kratzert et al., 2019a).

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3.2 Data

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Our experiments use the continental hydrology dataset, CAMELS (Catchment Attributes and Meteorology for Large Sample studies) (Addor et al., 2017). The CAMELS data set contains continuous meteorologic input, observed streamflow data, and catchment dependent spatially varying but temporally physical descriptors. CAMELS encompasses a total of 671 watersheds across the contiguous US. Due to some watershed delineation errors (Addor et al., 2017), we followed the suggestion from Kratzert et al. (2019a) to select 531 basins whose watershed boundaries are confirmed to be correctly delineated without digital errors. Each watershed is supplied with observed discharge and climate forcing data from remote sensing products (Daymet(Thornton et al., 2020), NLDAS(Xia et al., 2012), MAURER(Maurer et al., 2002)), climate models, and data assimilation with daily temporal resolution. Additionally, a corresponding hydrological model (SAC-SMA. Sacramento Soil Moisture Accounting model) is well calibrated for each watershed and its physical simulation is also available. Adopting such a wide distribution of watersheds, CAMELS provides a comprehensive and detailed physical description of watersheds. Selecting only a subset of those features as suggested by Kratzert et al. (2019a), we choose 27 physical descriptors from climatology, geomorphology and geology perspectives to characterize and discriminate across watersheds (Table A1 in the Appendix A.).

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These 27-d catchment physical descriptors are static vectors (\mathbf{x}^s) characterizing each catchment. We selected meteorological data from an updated version of MAURER as model dynamic input (\mathbf{x}^d), which are daily precipitation, daily minimum air temperature, daily maximum air temperature, average short-wave radiation, and vapor pressure. The observed discharge from USGS is our target variable (Q^O). Both daily meteorological weather inputs and discharge data cover a reasonably long record spanning from 1980 to 2014. The data for each catchment was partitioned into training and testing periods while building and evaluating deep learning models. Some experiments involved using a subset of training years or a subset of basins, therefore, we specify a default assessment scheme as to train a global model using data from 531 basins with 20

years of data. Under this assessment scheme, the training period starts from October 1st 1999 and ends on September 30th 2008. For a consistent evaluation, through all experiments, the testing period ranges between October 1st 1989 and September 30th 1999.

3.3 General setup

Among different LSTM-based models, We apply the same optimization algorithm (Adam optimizer (Kingma & Ba, 2017)) for training purposes to determine model parameters. Model parameters are learned from data and are thus continuously updated during training. The machine learning implementation also needs to specify hyper-parameters, which are set before training without learning from data. During training, hyper-parameters will not be updated. A few essential hyper-parameters include the look back period T and the dimension of hidden states $\mathbf{h}[t]$. Adopting the previous work's specification (Kratzert et al., 2019a) of these hyper-parameters, we determine T to be 270 days and the dimension of hidden states to be 256. For the details on other hyper-parameters (e.g., learning rate, batch size), please read the Appendix B in Kratzert et al.'s paper (2019a).

Machine learning models have uncertainties in model parameters after training. Initialized randomly, model parameters will often be optimized to different values during training. In simplistic terms, different model initializations will yield different models after training. Accounting for uncertainty, it has been shown that ensemble results from multiple model runs will facilitate the overall model performance (Kratzert et al., 2019a). Therefore, the streamflow prediction result in all following sections is an ensemble mean of five model realizations. For instance, the prediction of the EA-LSTM using physical descriptors is an average of five model predictions, which are optimized from different initializations. Note that for the Gaussian vector experiment, the randomness originates from two sources, including model initializations and the Gaussian vector assignment. For each of the five runs, their Gaussian vectors are assigned with different values.

Training deep learning models also requires a specification of the objective function. To account for cross-catchment variance, which is not considered in the commonly used mean squared error option, we use a smooth-joint NSE function (Kratzert et al., 2019a). The smooth-joint NSE function is shown below.

$$NSE^* = \frac{1}{B} \sum_{b=1}^B \sum_{t=1}^N \frac{(Q_t^m - Q_t^o)^2}{(s(b) + \epsilon)^2} \quad (15)$$

where B is the number of catchments, N is the number of daily data (days) for one catchment, which is indexed by b . Q_t^m is the predicted discharge at timestamp t ($1 \leq t \leq N$), while Q_t^o is the corresponding observed discharge. $s(b)$ is the standard deviation of the Q_t^o in basin b during training periods. ϵ is a constant term ($\epsilon = 0.1$) to avoid potential loss function explosion issue, which happens for catchments with extremely low $s(b)$.

For consistent model comparison, we're using the NSE score instead of RMSE (root mean squared error) to evaluate streamflow prediction. NSE is a metric suited particularly to evaluate hydrological predictions.

$$NSE = 1 - \frac{\sum_{t=1}^T (Q_t^m - Q_t^o)^2}{\sum_{t=1}^T (Q_t^o - \bar{Q}^o)^2} \quad (16)$$

Q^m is predicted discharge, Q^o is observed discharge, \bar{Q}^o is the mean of observed discharge. A NSE score of 1 indicates a perfect time series prediction.

3.4 State of the Art

In terms of data-driven regionalization methods, CT-LSTM and EA-LSTM have been shown to perform satisfactorily for the streamflow prediction task (Kratzert et al.,

2019a). To remind readers of the state-of-the-art performance which relies on the physical descriptors as shown in Table A1, in Figure 2, we show the testing NSE score for each catchment in the CAMELS dataset.

Table 1: State of the art LSTM based model. Mean and median refer to the summary statistics of the testing NSE scores across all 531 catchments in CAMELS.

Model	Mean	Median
Local LSTM	0.543	0.576
Global LSTM w/o static vectors	0.529	0.634
Global EA-lstm with 27-d descriptors	0.698	0.733

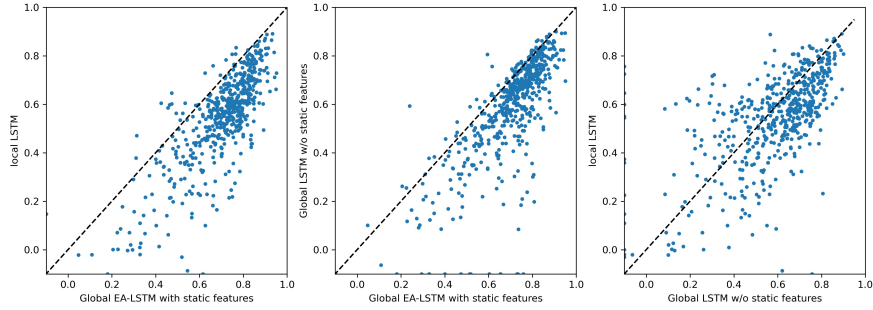


Figure 2: State of the art global regionalization performance using LSTM based deep learning architecture.

Local LSTM uses hydrologic data from only one catchment and does not need physical descriptors (\mathbf{x}^s) to combine data from multiple catchments. Thus, for 531 catchments, there are 531 Local LSTM models. On the other hand, global LSTM refers to a global model learned from the training data of 531 catchments. While the Global LSTM merges data from multiple catchments but does not use physical descriptors to adapt the network for different basins, the Global EA-LSTM with 27-d descriptors is also a global model trained and tested using all 531 catchments but it takes advantage of 27-d physical descriptors to perform robust regionalization. As shown in Table 1, both the mean and median of its NSE score is the highest (0.698 and 0.733 respectively) among the three model options. In this gauged prediction scenario, cross-catchment information sharing benefits global training and thus elevates predictive performance. These results have been previously shown by Kratzert et al. (2019a).

3.5 Proposed Approach

In this paper, our aim is to answer the question “How to perform regionalization when catchment physical descriptors are unavailable, uncertain, or of insufficient dimension?” To address this issue, we propose to assign a vector of random values as a surrogate for missing physical descriptors. Since a set of random vectors doesn’t have any similarity structure (i.e. correlation between any two random vectors is zero), they are a suitable baseline to incorporate the fact we don’t have any prior information on catchment similarity due to missing physical descriptors. By using these random vectors, we

enable the deep learning network to account for heterogeneity in catchment responses while sharing data across multiple basins.

Furthermore, the proposed concept of using random vectors as a baseline can also be used to evaluate the efficacy of known catchment physical descriptors. In other words, the performance difference between using random vectors and physical descriptors can imply the quality of physical descriptors. In section 4, we provide an extensive analysis of this concept in the context of streamflow prediction. In this paper, we consider two different strategies to create random vectors (Figure 3) as described below.

3.5.1 Gaussian Random Vectors



Figure 3: Random vector illustration. (a) refers to the d-dimensional Gaussian vector, while (b) illustrates the one-hot vector concept.

Figure 3a is a visual representation of the Gaussian vector (d-dimension) for all catchments. Random colors represent random numbers drawn from Gaussian distribution. In this strategy we assign d-dimensional vectors to each catchment where the vector values are drawn from a Gaussian distribution with zero mean and unit standard deviation. In other words, we randomly map each basin to a point in d-dimensional feature space.

3.5.2 One-hot Vectors

Figure 3b illustrates the one-hot vector representation. Each catchment is associated with a binary vector that is 1 for one dimension and is zero elsewhere. The dimension of the one-hot vectors equals the number of catchments. These one-hot vectors originated from the binary vectors used to encode categorical variables in regression, where in our case, the variable is catchment ID. There is one such one-hot binary vector for each basin and these vectors are orthogonal to each other. It bears emphasis that there's no freedom for the user to determine the dimension of the one-hot vector after the number of catchments in a global model is known. For k basins, the length of the one-hot vector for each basin is k. Although the one-hot vector does not involve random numbers, the randomness in this random vector assignment is from basin order. Regardless of how basins are sorted, one-hot vector assignment assures each basin will be assigned uniquely.

4 Experiments and Results

We evaluate the effectiveness of our random vector approach with a series of experiments. First, in Section 4.1 we compare the random vector performance to that of the state-of-the-art EA-LSTM model. Next, we investigate the applicability of the random vector approach under varying data richness scenarios. In Section 4.2, we create a data inadequacy scenario by limiting the number of basins used in the training data. We also examine the impact of limiting the number of years of training data, as demonstrated by the experiment in Section 4.3. To further assess the generalizability of the random

vector approach, we evaluate other model settings in Section 4.4 and present our analysis of the performance of the CT-LSTM model using random vectors along with the data inadequacy scenario. Additionally, we compare the efficacy of the EA-LSTM and CT-LSTM models using random vectors. In Section 4.5, we explore the practical implications of employing random vectors to model catchment complexities where physical descriptors for the system are incomplete. Finally, we show how the use of high-dimensional representation of catchments improves regionalization by distinguishing them from one another.

Implementing these experiments needs to specify a selective combination of the model architecture (EA-LSTM or CT-LSTM) and static vectors(\mathbf{x}^s). Options for \mathbf{x}^s include 27-d physical descriptors, random vectors, and mixing Gaussian vectors. For the simplicity of representing the results, we'll use acronyms to denote corresponding results of those experiments, that is, the combination of model architecture and \mathbf{x}^s . These acronyms are shown in the table 2. Models for the incomplete physical systems are not given acronyms. The comparisons across different models also involve statistical significance tests. Throughout the rest of this paper, we conducted pairwise statistical significance tests (Wilcoxon signed-rank tests) to evaluate model results differences. The statistical significance is evaluated using a 0.05 p-value threshold.

Table 2: This acronym table denotes the acronyms of model implementations. Combinations of model architecture and \mathbf{x}^s specifications are shown in their acronyms. The “d” in these notations represent the dimension of \mathbf{x}^s , which is only needed to specify the models using Gaussian vectors. For instance, EG-512 means EA-LSTM model using 512-d Gaussian vectors. ‘*’ means the corresponding models were not implemented.

\mathbf{x}^s		EA-LSTM	CT-LSTM
27-d physical descriptors		EP	CP
Random vectors	Gaussian d-dimension	EG-d	CG-d
	One-hot	EO	CO
Mixed Gaussian d-dimension vectors		EM-d	*

4.1 Effectiveness of Random vectors

To evaluate the applicability of our proposed random vectors method in regionalization, we first compared the predictive performance of a global model using random vectors (Gaussian or one-hot) against that using physically meaningful 27-d descriptors under EA-LSTM settings. The baseline model is the EP(Kratzert et al., 2019a) to show the state-of-the-art predictive performance. Substituting the 27-d physical descriptors with random vectors, our proposed method is implemented as either EG-d or EO.

Implementing Gaussian vectors requires a specification of d , which is determined empirically. The cumulative density function plot of the NSE score, shown in Figure 4 suggests using 512 (black solid line) as the Gaussian vector dimension because its testing performance is optimal compared to others.

The scatter plot (Figure 5) shows the testing NSE of the EG-512 and the EO versus the EP respectively across all 531 basins. Among these results, testing NSE scores less than -0.1 are forced to be -0.1 for illustration purposes. For each scatter plot, a cumulative density function (cdf) plot of NSE is also given. The EG-512 scatter plot is slightly upper skewed, the cdf of the EG-512 is also slightly right skewed compared to the EP. Figure 5a and Figure 5c shows that the EG-512 prediction performance is comparable

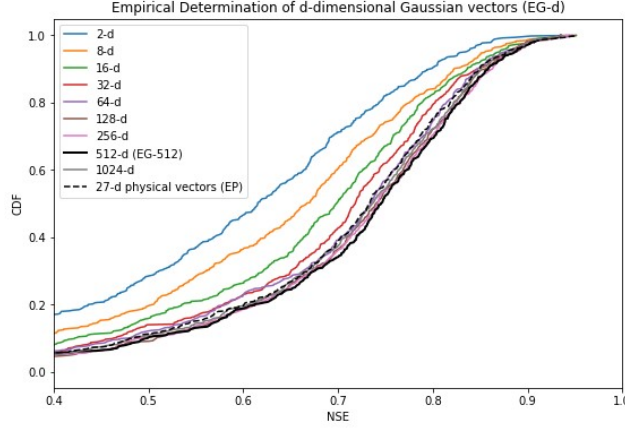


Figure 4: Cumulative density functions of the NSE score across different d Gaussian vectors for the EG-d. The X-axis is NSE score, which is truncated between 0.4 and 1 for a better illustration. The black dashed line represents the testing score corresponding to the EP. The black solid line corresponds to the EG-512, which yields the best performance in the EG-d.

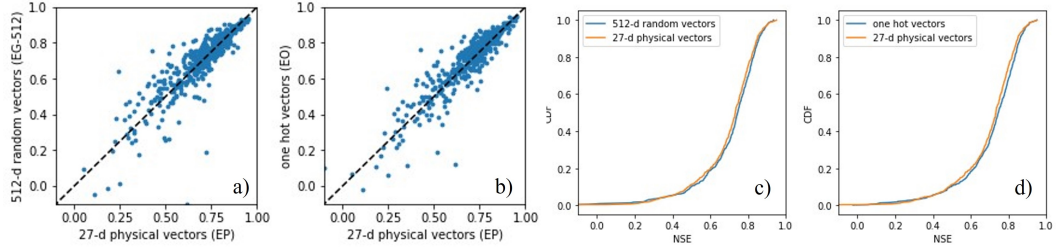


Figure 5: Performance comparison cross the EP, EG-512 and EO. Model architecture is EA-LSTM. (a) shows the NSE score scatter plot that compares EG-512 and EP while its cdf comparison is shown in (c). (b) shows the comparison between EO and EP while its cdf comparison figure is (d).

to, if not slightly better than, the EP (statistically significant). In Table 3, the mean and median of the EG-512 is 0.711 and 0.746, both of which yield slightly more satisfactory results than the EP. The same comparison between the EO and the EP also yields a sim-

Table 3: Performance comparison of the EA-LSTM using random vectors against physical descriptors. Statistical summaries across all 531 basins are in column ‘mean’ and ‘median’. The EG-512 and the EO are statistically different than the EP at 0.05 level.

Catchment static vectors	Mean	Median
27-d physical vectors (EP)	0.698	0.733
512-d random vectors (EG-512)	0.711	0.746
one-hot vectors (EO)	0.707	0.745

ilar trend. The mean and median of NSE score for the EO is 0.707 and 0.745. The EO reaches comparable prediction performance to the EP (statistically different at 0.05 level).

For the statistical comparison of Table 3 and all following tables (Table 4 to Table 10), their corresponding p-value can be found in the supplemental information (Table S1 to Table S9).

As we can see, using random vectors gives performance comparable to using known physical descriptors. Furthermore, the random vector approach leads to significantly better results when compared to other strategies that do not use known physical descriptors (i.e. Figure 2, building local models or trivial merging of data from multiple basins). Hence, the random vector approach is a viable solution when catchment characteristics are not available. This performance is evaluated using the standard setting (section 3.2, 10 years training data from 531 basins). Although such abundant training data shows slightly elevated testing performance, the proposed random vector method might still be inapplicable in data poor situations. To assess the impact of data sparsity, we conducted an exhaustive analysis on the different data inadequacy scenarios with either fewer number of basins or fewer number of training years.

4.2 Effect of number of basins

For this situation we’re creating a data inadequacy scenario where the training data consists of a limited set of k basins. Such a group of limited basins forms an insufficient global hydrologic dataset to train an LSTM based model. This experiment aims to answer the question ”Given only k basins without physical descriptors, will the proposed random vector strategy be applicable for regionalization?” We vary k from 10 to 50 to 100 and follow the default assessment scheme as outlined in Section 3.2.

To generate the basin sets, we randomly select k basins as a group repetitively without replacement until all basins are selected. When the remaining basins cannot form a group with exactly the size k , those basins are either merged with the last group or form a stand-alone group as long as its order of magnitude approximates to k . For instance, when selecting 10-basin group, we select 53 groups in total, and the last group contains 11 basins. Similarly, the last group (11th group) in the 50-basin group has 31 basins. The last group (fifth group) in 100-basin group has 131 basins.

For the 53 groups of 10-basin groups, we compare the predictive performance using random vectors relative to the performance of the model using 27-d physical descriptor. This comparison is illustrated in Figure 6. The X-axis denotes one-hot vector and Gaussian vectors (varying d). Each category shows a box plot of performance comparison across basins. Median (blue dots), 25th percentile and 75th percentiles (upper and lower box line) are shown for each box. Black hollow circles outside the upper and lower box lines are outliers outside the specified quantile range. The Y-axis is the NSE score improvement for each individual catchment compared to the 27-d physical descriptors. The red line indicates the threshold for improved performance. A box plot whose NSE distribution is skewed to positive NSE score improvement (above the threshold line) indicates a general performance improvement in that random vector category. Both the EG-256 and the EG-512 show an performance improvement more pronounced than other Gaussian vector dimensions and one-hot vectors. They both improve the NSE score on an average of 0.082 (or in median 0.066). The reference performance of the EP is reported at the table 4. The mean predicting NSE score is 0.308 while the median is 0.317, both show that the 10-basin group downgrades the model performance because fewer basins provide only limited training data and thus constrains model learning generalizable hydrologic behavior.

For the 50-basin group and 100-basin group, the plot of NSE improvement is shown in Figure 7 except that we plot only the median of each case for a succinct visualization. The red line also marks the performance improvement threshold. Table 4 summarizes the NSE score improvement for all cases. As data from a greater number of basins are involved, the model performance gradually increases from 0.317 (10-basin) to 0.599 (50-

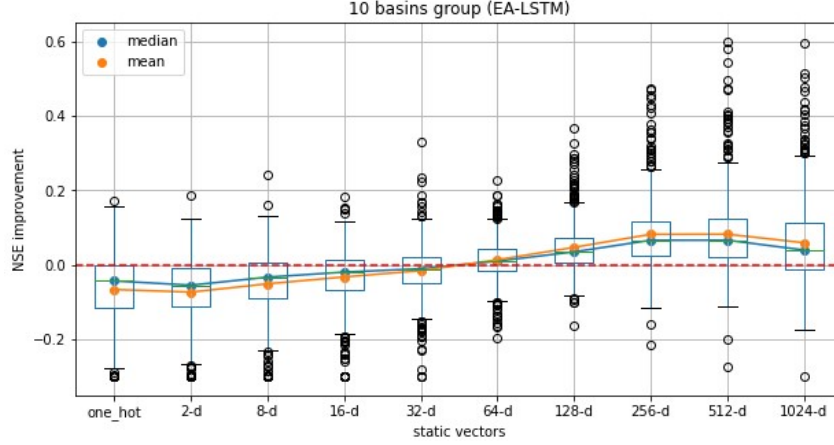


Figure 6: Random vectors implementation for 10-basin group EA-LSTM. Categories along the X-axis represent random vectors, including one-hot vectors (length of 10) and Gaussian vectors (dimension d varies from 2 to 1024). The Y-axis shows the individual basin NSE score improvement of the random vector in contrast to its corresponding EP, which is trained using the same basins. A zero NSE improvement indicates an improvement threshold marked by the red line. Within each category, 531 NSE improvement scores are distributed in the box plot where outliers exceeding 25th and 75th quantile are marked by black hollow circles.

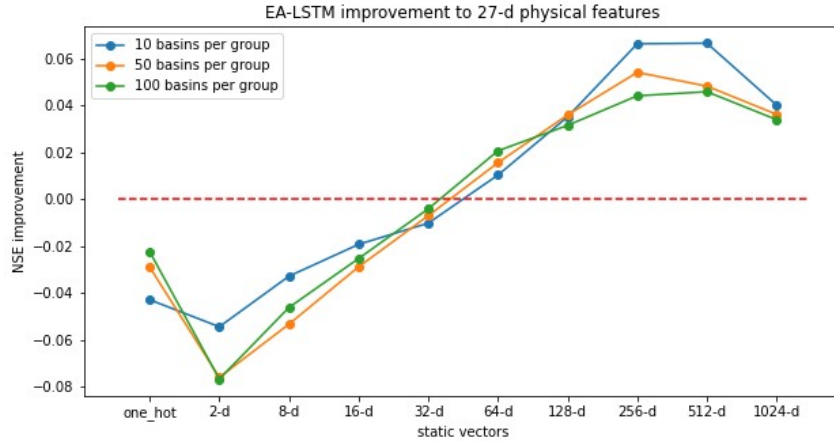


Figure 7: Random vectors implementation for k -basin group EA-LSTM. k varies from 10 to 50 to 100. The median in Figure 6 are blue lines. Within each random vector category as shown in X-axis, the median of the individual-basin NSE score improvement in contrast to EP for k basins is plotted. Orange dots are the 50-basin group while green color represents 100-basin group

basin) to 0.656 (100-basin), all of which are lower than 0.733, the performance of the model using all 531 basins. Note that both the trend and the performance are comparable to the previous work, where the impact of the training data inadequacy on the EA-LSTM performance is explored (Gauch et al., 2021). Table 4 and Figure 6 show a consistent performance improvement comparison. Regardless of how limited the number of basins, the Gaussian vector strategy (with an optimal dimension of either 256 or 512) slightly improves the 27-d physical vectors. In particular, the performance improvement from the Gaussian vectors becomes saturated when d reaches 256 or 512. For 50-basin group,

Table 4: The individual-basin performance improvement of the EG-d and the EO to the EP. Note that the EP row does not show the NSE improvement, instead it shows the NSE score performance, which is the red dashed line performance in Figure 6 and 7. The largest performance improvement as indicated by the positive largest numbers is in bold font.

k-basin group		10	50	100	10	50	100
d		mean			median		
Gaussian vector (EG-d)	2	-0.073	-0.085	-0.09	-0.054	-0.076	-0.077
	8	-0.051	-0.061	-0.058	-0.033	-0.053	-0.046
	16	-0.032	-0.031	-0.03	-0.019	-0.029	-0.025
	32	-0.015	-0.01	-0.008	-0.01	-0.007	-0.004
	64	0.014	0.018	0.025	0.01	0.016	0.021
	128	0.047	0.045	0.039	0.035	0.036	0.031
	256	0.082	0.062	0.053	0.066	0.054	0.044
	512	0.082	0.055	0.053	0.066	0.048	0.046
	1024	0.06	0.038	0.039	0.04	0.036	0.034
one-hot (EO)		-0.066	-0.035	-0.03	-0.043	-0.029	-0.022
27-d physical descriptors (EP)		0.308	0.569	0.620	0.317	0.599	0.656

the average of the single-basin NSE improvement is 0.062 at 256-d while the median of the single-basin NSE improvement is 0.54. For 100-basin group, EG-512 improves the NSE score slightly with a mean of 0.053 while the median is 0.046, which is approximately the same to the extent of what EG-256 improves. When the dimension of the Gaussian vector becomes a higher 1024-d, the performance improvement begins to degrade as indicated by a smaller NSE improvement. In summary, we show that random vector approach shows robust performance even with fewer number of catchments in the dataset and hence can be used in situations where only few catchments are available.

4.3 Effect of number of training years

In addition to the number of basins, another perspective on data inadequacy is the number of training years. Varying the training years from 1 to 2 to 5 years, we sought the answer to this question “Given only a few years of training data, will the proposed random vector strategy be applicable for regionalization? ” An LSTM model is trained for all 531 basins with a limited number of years. The EG-d is tested against the EP under three sparse data cases, which are 1 year of data (October 1st 2007 to September 30th 2008), 2 years of data (October 1st 2006 to September 30th 2008) and 5 years of data (October 1st 2003 to September 30th 2008). Models are tested for the same years (October 1st 1989 to September 1st 1999) for consistent comparison.

Our previous empirical analysis indicates an optimal specification of d (Gaussian vector dimension) to be 512 (Section 4.1), so the implementation of basin random vectors includes either 512-d Gaussian vectors or one-hot vectors. The Figure 8 shows that both random vector strategies lead to prediction performance similar to the case utilizing 27-d physical descriptors. For the reference, as the number of years of training data increases, the performance of EP also increases from 0.632 (1 year) to 0.697 (two years) to 0.766 (five years). This increasing trend was also identical to what Gauch et al. showed. In particular, the EG-512 yields a more satisfactory performance than the EO. As shown in Table 5, a NSE score improvement (both in mean and median) is observed when implementing 512-d Gaussian vectors, while the NSE score improvement is only observed when using 5 years of training data when the one-hot vector strategy is applied. The re-

sults show that even when training data are limited, randomly assigned vectors are still able to learn as well as 27-d physical features.

Table 5: The impact of the number of training years on the performance improvement of random vectors for EA-LSTM. “Mean” and “Median” refer to statistics of the individual-basin NSE score improvement in relative to EP. The EP row does not show NSE score improvement, instead it shows the NSE score performance, which is the reference performance in Figure 8. Positive numbers mean that random vectors yield better predictive performance.

Number of training years		1	2	5
Gaussian 512-d (EG-512)	mean	0.013	0.052	0.026
	median	0.009	0.041	0.019
one-hot (EO)	mean	-0.025	-0.003	0.015
	median	-0.023	-0.005	0.013
27-d physical descriptors (EP)	mean	0.399	0.628	0.719
	median	0.632	0.697	0.766

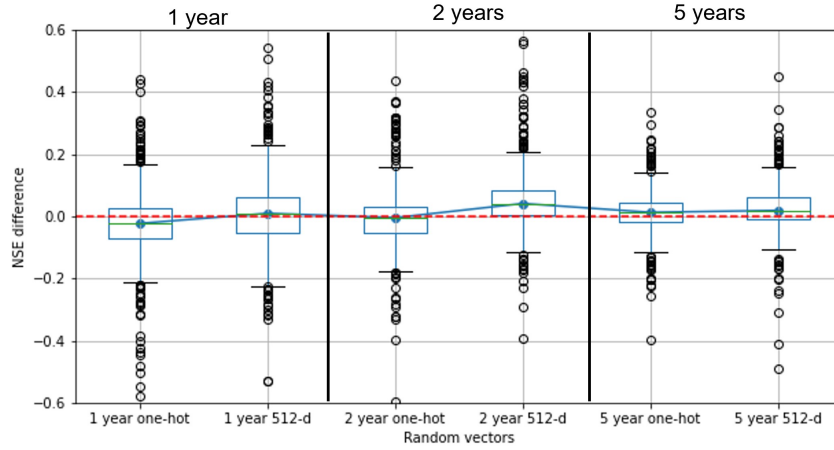


Figure 8: The impact of the number of training years on EA-LSTM. The Y-axis represents the individual-basin NSE score difference between the corresponding category in X-axis and the predictive performance using 27-d physical descriptors (EP). The red line indicates performance improvement threshold.

4.4 Performance of alternative models

As outlined in the section 3.1, both EA-LSTM and CT-LSTM adopt \mathbf{x}^s in different ways. From previous sections (section 4.1, 4.2, and 4.3), we’ve shown the efficacy of the random vectors in EA-LSTM in both data rich and data poor scenarios. It remains unknown whether random vectors are applicable to the CT-LSTM. Experiments of this section are designed to clarify this doubt.

We first evaluated the performance of random vectors under the CT-LSTM setting to answer this question “Under different model architectures, will the proposed random vector strategies be applicable for regionalization?”. We then examined the region-

alization performance of random vectors across the EA-LSTM and the CT-LSTM to answer the question “Which random vector strategy is better suited for regionalization, Gaussian vectors or one-hot vectors?” Last, we selected the CT-LSTM as the model architecture for an exhaustive analysis on the data inadequacy cases in terms of basin numbers.

4.4.1 Random vectors in the CT-LSTM

For the CT-LSTM, the Gaussian vector implementation needs to specify the optimal vector dimension d . Figure 9 shows that the CG-16 yields the most satisfactory performance among different Gaussian vector dimension options. Therefore, we empirically select 16 as the optimal Gaussian dimension to represent the CG- d performance (Figure 10c). Note that the optimal 16- d of the CG- d is less than the optimal 512- d of the EG- d . We’ll explain this in the section 5.1 in “Discussion” section. Using 27- d physical descriptors, CP achieves performance comparable and slightly better than EP (Figure 10a and Table 6). The median NSE score performance improves from 0.733 (CP) to 0.744 (CO). Random vector options (CO and CG-16) slightly outperform 27- d physical descriptors (CP). The median of testing NSE performance improves from 0.744 to 0.754 when using the one-hot vector strategy, while the CG-16 elevates the performance to 0.752.

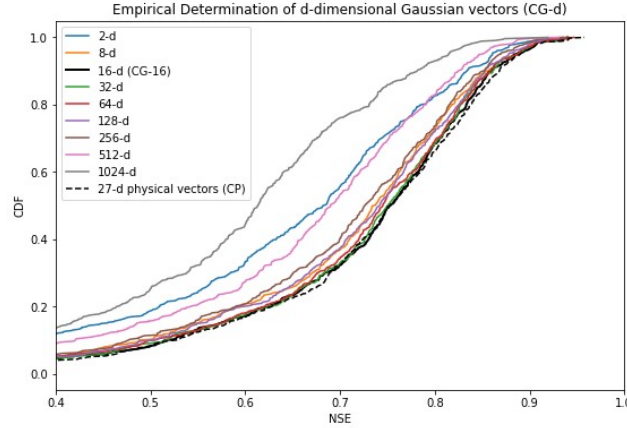


Figure 9: Cumulative density function plots of the NSE score across different d Gaussian vectors for the CT-LSTM. The X-axis is truncated between 0.4 and 1 for a better illustration. The black dashed line represents the testing score of the CP, the black solid line corresponds to the optimal 16- d performance among the Gaussian vector groups (CG-16).

Table 6: Random vector comparison cross different models (The CT-LSTM based random vector performance is statistically different from the CP at 0.05 level).

models	Mean	Median
EP	0.698	0.733
CP	0.715	0.744
CO	0.720	0.754
CG-16	0.717	0.752

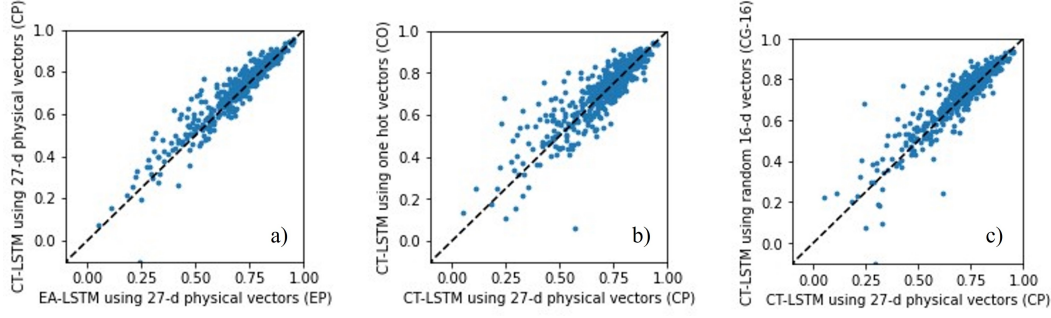


Figure 10: Predicted performance comparison of a random vector implementation in CT-LSTM (CO and CG-16) in contrast to CT-LSTM using 27-d physical descriptors (CP). (a) is the comparison between CP and EP; (b) is the comparison between CO and CP; (c) is the comparison between CG-16 and CP.

Although the slight improvement of the CO and the CG-16 in contrast to the CP imply the applicability of random vectors in the CT-LSTM, data abundance has always been an important factor impacting the machine learning model performance. To consolidate the argument that CT-LSTM with random vectors, especially one-hot vectors, yields better performance consistently under various data richness scenarios, we repeated the experiments outlined in section 4.2 for the CT-LSTM. Training data are limited by the number of basins.

Table 7: The improvement of random vectors over 27-d physical features in the CT-LSTM. “Mean” and “Median” refer to statistics of NSE score improvement for individual basins in relative to the CP. Note that the CP row shows the NSE value for the CP, instead it shows the NSE score performance, which is the reference performance in Figure 11 and 12. The most satisfactory performance is in bold font: 32-d Gaussian vector, 64-d Gaussian vector, and one-hot vector. For 10-basin and 50-basin group, their NSE performance difference between random vectors and the CP counterpart is significantly different at 0.05 level, while the NSE difference comparison at 100-basin group does not show statistical significance.

k-basin group		10	50	100	10	50	100
	d	mean			median		
Gaussian vector (CG-d)	2	-0.079	-0.073	-0.074	-0.075	-0.063	-0.063
	8	-0.055	-0.031	-0.024	-0.050	-0.028	-0.023
	16	-0.037	-0.015	-0.007	-0.037	-0.018	-0.010
	32	-0.022	-0.006	0.004	-0.023	-0.008	0.001
	64	-0.023	-0.004	0.002	-0.021	-0.006	0.000
	128	-0.041	-0.019	-0.019	-0.039	-0.016	-0.003
	256	-0.074	-0.059	-0.033	-0.072	-0.048	-0.026
	512	-0.103	-0.125	-0.094	-0.097	-0.114	-0.083
	1024	-0.134	-0.188	-0.174	-0.129	-0.191	-0.171
one-hot (CO)		-0.046	-0.007	-0.005	-0.042	-0.005	0.001
27-d physical descriptors (CP)		0.454	0.655	0.684	0.481	0.687	0.709

Figure 11 exhibits a box plot showing the NSE improvement for the 10-basin group using the CT-LSTM architecture. Any point above the red line (NSE score improvement threshold) indicates a performance improvement in contrast to 27-d physical descriptors.

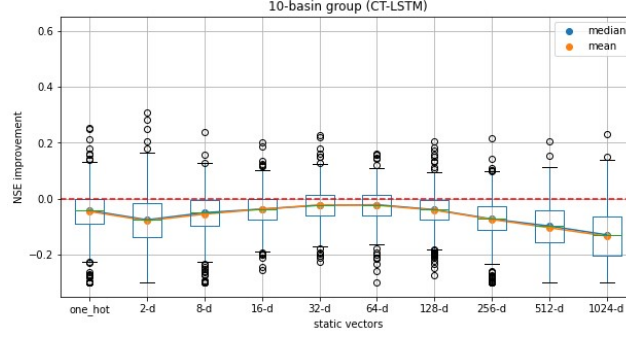


Figure 11: Impacts of random vectors on CT-LSTM for a 10-basin group. Categories on the X-axis represent random vectors, including one-hot vectors (length of 10) and Gaussian vectors (dimension d varies from 2 to 1024). The Y-axis show the NSE score improvement for individual basin of the random vectors in contrast to the CP. A zero NSE improvement indicates no performance improvement marked by the red line.

In the 10-basin group category, the optimal Gaussian d for the CG-d is lower than that of the EG-d. The optimal Gaussian vectors performance is comparable to that of one-hot vectors. To obtain a general insight, we varied k from 10 to 50 to 100 and therefore produced the following result in Figure 12 and Table 7.

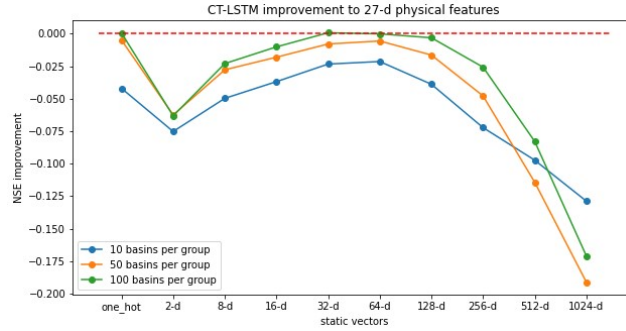


Figure 12: A random vector implementation for a k -basin group CT-LSTM. k varies from 10 to 50 to 100. Within each random vector category as shown in X-axis, the median of NSE improvement score for individual basin in contrast to the CP for k basins is plotted. Blue dots are 10-basin group, orange dots are 50-basin group, while green color represents 100-basin group

Figure 12 shows the median of the NSE improvement using random vectors in the CT-LSTM in contrast to the CP. Dots below the red line mean the prediction performance of the corresponding categories is worse than the CP. As the number of catchments available for training increases, the one-hot vector strategy and optimal Gaussian vectors in the CT-LSTM yields performance comparable to the CP. The optimal d for the CG-d is either 32 or 64, which is lower than the optimal 512-d in the EG-d. As also recognized in Figure 10, this discrepancy of optimal Gaussian d between the CT-LSTM and EA-LSTM can be explained by the number of parameters involved in these model architectures and we'll expand this discussion in section 5.1. We point out that these random vector strategies are approximate to but do not marginally exceed the CP performance. In particular, the relative significant performance improvement occurs when us-

ing the one-hot vector in the 100-basin group but to a much lesser extent. Varying k from 10 to 50 to 100, as more catchments are involved until 531 basins are included, the one-hot vector is a preferable random vector strategy for CT-LSTM than Gaussian vectors.

4.4.2 Best performance of using random vectors

In the CT-LSTM setting, the above experiments demonstrate that random vector strategies still prevail over 27-d physical vectors. The best random vector strategy for the CT-LSTM is one-hot vector, while the best random vector method for the EA-LSTM is 512-d Gaussian vectors. The preferable random vector strategy varies depending on the model setting.

In a pursuit of model performance when utilizing random vectors, we need to provide a practical solution to the question “When implementing random vectors to perform regionalization, shall I use Gaussian vectors or one-hot vectors?”. We next compared the optimal random vector performance between the CT-LSTM and EA-LSTM. Figure 13 shows the testing NSE difference from the various EG-d against the CO. Based on the previous result showing that the EO is not as good as EG-d, ‘one-hot’ on X-axis (EA-LSTM random vector strategy) is omitted. Figure 13 shows the median of the NSE difference for various selections of k basins. All points are below the performance threshold line, indicating that the CO slightly outperforms EG-d. When implementing the random vector strategy as a surrogate for missing physical descriptors, the best performance is obtained when applying CO.

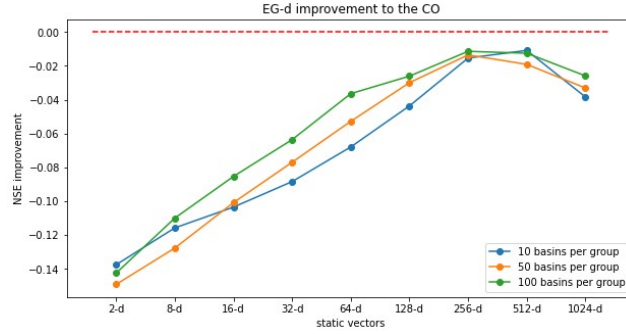


Figure 13: The performance of EG-d in contrast to the CO for k -basin group. k varies from 10 to 50 to 100. The Y-axis is the NSE difference score quantifying the performance of the EG-d (categories in X-axis) relative to the CO for individual basin. The red line marks no NSE difference. Plotted points are median of the NSE difference across all basins. For points below the red line, they mean that the CO yields more satisfactory performance

4.5 Incompleteness of physical descriptors

So far we considered the scenario where physical descriptors are not available and assessed the performance of our random vector approach. In this section, we consider a more common regionalization challenge where physical descriptors are incomplete. Incomplete physical descriptors under-represent a system of catchments and can only help regionalization in a limited degree. To tackle this problem, the question becomes “will the proposed random vector strategy benefit the model regionalization in this information deficient physical system?” To assess the performance caused by this deficiency, we define a physically underrepresented global system in CAMELS where only a subset of 27-d physical descriptors is used to distinguish basins. We compare the global LSTM us-

ing random vectors in contrast to the global model using these insufficiently informative descriptors. One extreme case is a system without any static catchment descriptors, which has been shown in the section 3.4 (Figure 2).

Ignoring the model selection differences, we select EA-LSTM for this experiment because it explicitly modulates LSTM via static vectors. The EA-LSTM using some subset of 27-d physical descriptors is trained and compared. EA-LSTM using 9-d climate features, 10-d geology features, and 8-d geomorphology features are trained separately and compared to the EA-LSTM using random vectors.

Catchment hydrologic models are formulated to resolve complexity and associated scaling issues in hydrological processes. Both issues will require a comprehensive physical understanding. From a practical perspective, static physical descriptors (for instance, Table A1) can only characterize complex catchments to a limited dimension because a sufficient catchment complexity characterization is challenging across scales. In the field scale, a hydrological model might characterize local hydrological processes completely, but the applicability of this locally built model to a larger basin might fail if the model is not adjusted, either simplified (reduce the number of parameters) or made more complex (enrich physical parameters). Therefore, for the regionalization involving catchments at various scales, the question becomes “Are any given physical descriptors sufficient for modeling the complexity of catchments?” This question also implies another question: “how many physical dimensions do we need for characterizing the complexities of stream-flow generation processes?”

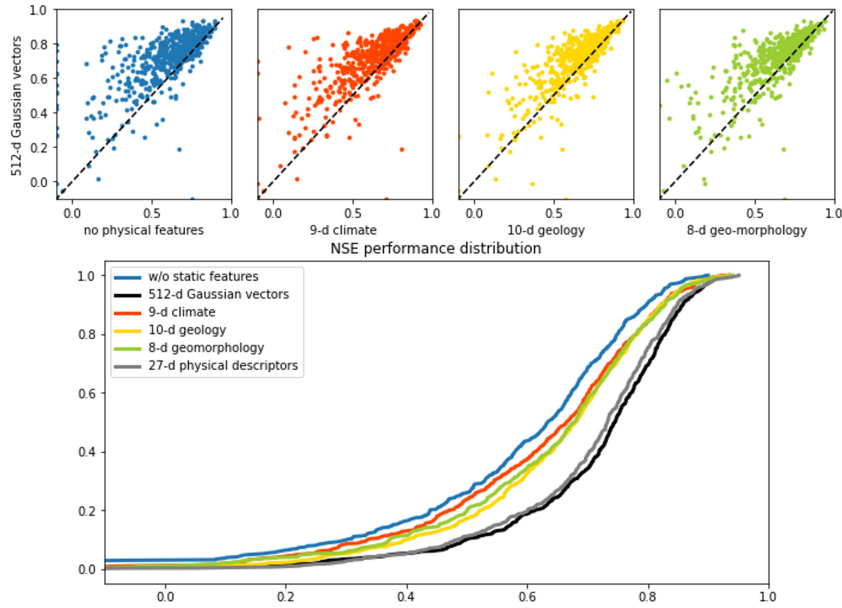


Figure 14: The performance of the EG-512 in contrast to EA-LSTM under a physically incomplete catchment system. Scatter plots show testing NSE scores comparison between the Y-axis and the X-axis. The y label is EG-512 and is fixed in the above 4 figures. The X-axis changes from a no physical feature system, a climate feature system, a geology feature system, to a geo-morphology feature system. The below cumulative density function plot collects NSE scores together for each category aforementioned. The black line is EG-512. We also plotted the benchmark performance with sufficient 27-d physical descriptors (grey solid line, EP) to remind the reader of the performance under a physically sufficient catchment system.

Table 8: Physical system completeness identification. The most satisfactory performance is in bold font (EG-512). In contrast to the EP, EG-512 and other physically incomplete system shows statistically different performance (at 0.05 level).

catchment static vectors	Mean	Median
(0-d) Without static features	0.529	0.634
512-d Gaussian vectors (EG-512)	0.707	0.745
9-d climate features	0.611	0.665
10-d geology features	0.638	0.679
8-d geomorphology features	0.630	0.680
27-d physical descriptors (EP)	0.698	0.733

To answer these questions, we compared our random vector results to models utilizing incomplete sets of physical vectors. In Table A1, 27-d physical descriptors are categorized into three groups: climate, geology and geo-morphology. Among these, the descriptors of any single group are an under-representative description for basins. For instance, 9-d climate descriptors presumably characterize basins less informatively than 27-d physical descriptors. For this experiment, we choose the EA-LSTM as the model structure and use 512-d Gaussian vector as its optimal random vector strategy. Each one of the three descriptor subset groups leads to an EA-LSTM under a physically uninformative system since complexities are simplified and the system incurs information loss. For the extreme case where there are no physical descriptors present, the global model is a simple global LSTM without basin characteristics, results of which were shown early in section 3.4 (Figure 2).

In Figure 14, a distribution of scatters above the diagonal line (exactly equal performance from the methods indicated by axes) indicates that Gaussian 512-d vectors outperform all these physically incomplete conditions. This fact is better illustrated in the cumulative density function plot as the distribution of NSE scores is skewed to upper tail. Both its mean and median NSE scores are higher than any physically incomplete characterization (Table 8). Note that as shown earlier, the EG-512 case reaches comparable and slightly better performance than EP. This observation also implies that 27-d physical vectors are lacking additional physical characterizations.

4.6 Effectiveness of Distinguishing Basins in the High-dimensional Space

Missing or incomplete physical descriptors make catchments less distinguishable from each other. Even for the assumed complete 27-d physical descriptor, they suffer from losing information as heterogeneous catchment systems are spatially simplified. Of the 27-d physical descriptors, the spatially dependent descriptors are deterministic representations of catchments, such as soil porosity, silt fraction, etc. This simplification also reduces the functionality of static vectors to distinguish catchments from each other, which might produce disadvantages for regionalization.

Further, the proposed random vector strategy projects catchments in the high-dimensional space. In particular, the EG-512 assigns a Gaussian vector of 512-dimension to catchments, while the CO uses the 531-d one-hot encoded vector to represent catchments. In other words, it seems that characterizing catchments in a high-dimensional space distinguishes them from each other and thus improves regionalization. Recognizing this, the question becomes: “Can we incorporate 27-d physical descriptors in the high-dimensional space?”

We offer two methods to include the 27-d physical descriptors in the high dimensional space and compare the performance of these methods with the performance of the random vector approach. We use EA-LSTM in these experiments rather than CT-LSTM because training CT-LSTM increases the computational burden and complicates machine learning. Additionally, EA-LSTM explicitly modulates the LSTM architecture. In the first method, we concatenate the 27-d physical descriptors with additional Gaussian vectors to expand the dimension of \mathbf{x}^s . We refer to this as the mixed Gaussian vector approach. In the second method, we create an embedding layer to explicitly project \mathbf{x}^s into a high dimensional space before entering the EA-LSTM cell.

4.6.1 Mixed Gaussian Vector

The 27-d physical vectors are augmented with extra dimensions filled by Gaussian vectors, which is named as “mixed Gaussian vectors” (denoted as ‘EM’). Catchments are gradually more distinguishable as their Gaussian vector dimension increases. These appended Gaussian vectors improve the distinctiveness of catchment characterization. We define a global system with 64-d, 128-d, 256-d, and 512-d vectors, all of which include the 27-d physical descriptors. For instance, for the 64-d \mathbf{x}^s , besides the 27-d physical descriptors, 37-d (64-27=37) vectors are randomly drawn from the Gaussian distribution.

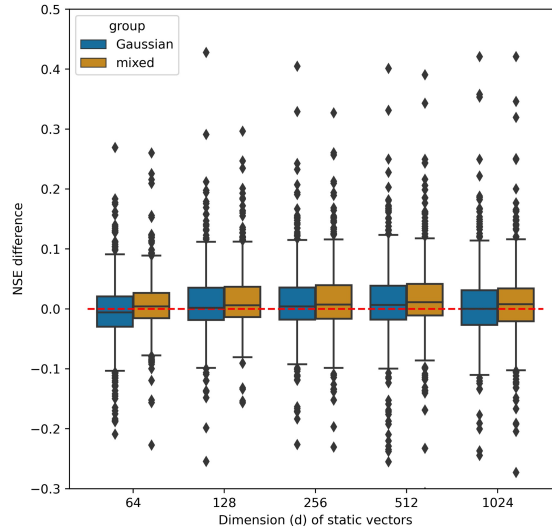


Figure 15: Comparison of the performance between Gaussian vectors (EG-d, blue box) and mixed Gaussian vectors (EM-d, flaxen box). The X-axis is the dimension of the static vector (from 64 to 1024), while the Y-axis shows the NSE difference in contrast to the EP for individual basin. The red line specifies the performance improvement threshold. Box portions above the red line indicate performance improvement.

As shown in Figure 15, compared to the baseline performance, which is the EP, the mixed Gaussian vector (EM-d) yields better performance and achieves the maximal performance improvement at EM-512. On average, the NSE improvement is 0.018. From 64-d, to 1024-d, all of the EM-d results yield better performance (positive NSE score improvement statistics in Table 9). In contrast to a pure random Gaussian system, given the same d Gaussian dimension (blue and flaxen box in the same X-axis category), the

Table 9: NSE performance difference of the mixed Gaussian vectors (EM-d) and Gaussian vectors (EG-d) in contrast to 27-d physical vectors (EP) for individual basin. Positive scores mean that the EP yielded worse predictive performance (these results are statistically different from the EP at 0.05 level). The highest value is highlighted in bold.

static vector dimension (d)		64	128	256	512	1024
Gaussian vectors (EG-d)	mean	-0.007	0.010	0.010	0.009	0.002
	median	-0.006	0.002	0.004	0.007	0.000
Mixed Gaussian vectors (EM-d)	mean	0.009	0.013	0.014	0.018	0.009
	median	0.004	0.006	0.007	0.011	0.008

EM-d also marginally improves the NSE score. In Table 9, the NSE improvement in ‘Mixed Gaussian vectors’ are consistently more pronounced than ‘Gaussian vectors’ across varying static vector dimension d . The mixed Gaussian vector leads to marginally better global model performance compared to either pure random Gaussian vector system or pure physical system. As such, it suggests that when physical descriptors are augmented with Gaussian vector in a high dimension space, catchments are more distinctively represented, which supports and benefits regionalization.

4.6.2 Additional Embedding

Besides the mixed Gaussian vector approach, the other method to characterize catchments in a high dimensional space is to create an embedding layer between x^s and the input gate i of the LSTM cell. That is to say, the equation 9 in EA-LSTM is replaced by 17 and 18:

$$\mathbf{v} = \sigma(\mathbf{W}_v \mathbf{x}^s + \mathbf{b}_v) \quad (17)$$

$$\mathbf{i} = \sigma(\mathbf{W}_i \mathbf{v} + \mathbf{b}_i) \quad (18)$$

where \mathbf{v} is the embedding layer that x^s is mapped into. We chose 512 as the dimension of \mathbf{v} for its empirical outstanding performance in the EG-512. We first mapped the physical descriptors into a 512-d embedding layer (denoted as ‘PEA’). Then as its random counterpart when \mathbf{x}^s is not available, we also experimented to map random vector into the embedding layer (denoted as ‘REA’). To preserve the modeling capacity without including additional model parameters, the dimension of the random vectors in the REA is still 27.

Table 10: Performance comparison of the EA-LSTM using additional embedding layers against previous random vector approaches. Statistical summaries across all 531 basins are in column ‘mean’ and ‘median’. This result is statistically different from the EP at 0.05 level.

Catchment static vectors	Mean	Median
27-d physical descriptors (EP)	0.698	0.733
512-d Gaussian vectors (EG-512)	0.711	0.746
27-d physical descriptors with 512-d embedding (PEA)	0.713	0.753
27-d Gaussian vectors with 512-d embedding (REA)	0.714	0.757

As shown in Table 10, the mean of testing NSE scores for the ‘PEA’ is 0.713 while its median is 0.753. With similar performance, the mean and the median of the testing NSE scores for the ‘REA’ is 0.714 and 0.757 respectively. Note that both ‘PEA’ and ‘REA’ shows slightly better performance than EG-512. In particular, ‘PEA’ and ‘REA’ yields comparable performance.

5 Discussion

This section is organized as four subsections. The first section (5.1) presents comparison across the experiments in Section 4.1, 4.2, 4.3, 4.4 and shows the presence of an optimal large d . The second section (5.2) compares the results in Section 4.5 and 4.6 and discusses the regionalization advantage from high-dimensional characterization of catchments. The third section (5.3) presents the analysis of the embedding layers in both EG-512 and EP. It also shows the discussion on what all of our results will imply to understanding catchment similarities and complexities. The forth section (5.4) presents a discussion on the impact of our results towards the understanding of deep learning in the context of streamflow prediction, and shows one practical utility of random vectors in assessing the completeness of physical descriptors. The fifth section (5.5) outlines the limitations of the current study and describes possible future directions.

5.1 Random Vectors

Results in section 4.1 show that the proposed random vector method achieves a performance comparable to the state-of-the-art model (Figure 5, Table 3). In other words, without any knowledge of physical descriptors, the global LSTM based model using random vectors successfully learns universal hydrologic behavior and sustains benchmark streamflow prediction performance. These random vectors retain practical feasibility without having to obtain any physical descriptions of basins. This is arguably the most significant scientific contribution of this paper.

The exhaustive analysis from section 4.2, 4.3, and part of 4.4 verifies the applicability of employing random vectors in data scarce regions. For a limited number of basins (Figure 6, 7, 11, 12, Table 4, 7) and a few years of training data (Figure 8), the two situations which restrict hydrologic extrapolation across catchments, random vectors are still viable for hydrologic regionalization.

The prediction performance achieved by random vectors varies between the EA-LSTM and the CT-LSTM, which we hypothesize is the result of different modulation levels in their architectures. Random vectors functionalize as static vectors (\mathbf{x}^s) that represent each basin, from which the LSTM family models modulate its internal computation and mapping across neurons for each basin distinctively. That is, for a given weather input \mathbf{x}^d , the global model is aware of which basin the \mathbf{x}^d data originates from and thus modulates how streamflow shall be predicted differently in contrast to other basins. Yet this modulation extent likely varies between EA-LSTM and CT-LSTM. CT-LSTM concatenates \mathbf{x}^s with \mathbf{x}^d at each timestamp and thus performs a stronger modulation because this catchment awareness is passed through all gates in the LSTM. Merely feeding into the input gate, the EA-LSTM does not modulate the network as well as the CT-LSTM but it ensures \mathbf{x}^s is involved in the temporal context update (memorizing and forgetting). Across those different LSTM model selections with various catchment modulation degree, random vectors consistently perform as well as, if not better than, physical descriptors for learning across basins (Table 3, 6).

Random vectors can either be Gaussian vector or one-hot vector. For the Gaussian vector, note that its implementation needs to specify its dimension d , which is empirically obtained. The optimal d for the Gaussian vector is different between the EA-LSTM and the CT-LSTM. For the EA-LSTM, the optimal d is either 256 or 512 (Fig-

ure 4, 7, 6), while for the CT-LSTM, it is in the range of 16 to 64 (Figure 9, 11, 12). This means that the EA-LSTM needs a higher dimension of static vectors to perform regionalization than the CT-LSTM. We explain this difference by the amount of trainable machine learning parameters. The increasing number of trainable model parameters of CT-LSTM hinders the training processes. For the CT-LSTM, an increased static input will expand the concatenated input $\mathbf{x}[t]$ dimension (Equation 8), which in turn enlarges the dimension of the transformation matrices \mathbf{W}_i , \mathbf{W}_f , \mathbf{W}_g , \mathbf{W}_o . In contrast, the static input (\mathbf{x}^s) dimension only impacts EA-LSTM's input gate dimension (\mathbf{W}_i). Consequently, given the same \mathbf{x}^s dimension augmentation, the parameter increment of CT-LSTM is four times the increase of the number of parameters in EA-LSTM. A higher d-dimension Gaussian vector CT-LSTM becomes more difficult to optimize than that for the EA-LSTM.

Although the optimal d differs between the CT-LSTM and the EA-LSTM, the performance saturation trend is identical. As illustrated by the results between section 4.1 and 4.4, when expanding static vector dimension, the predictive performance saturates at a certain point and then deteriorates. This pattern indicates a presence of the optimal d , which cannot be too large or too small and is suitable for achieving best model performance. In particular, the optimal d is universal regardless of the number of basins involved (Figure 6, 7, 11, 12) as well as the number of training years (Figure 8). Thus, it suggests implications for addressing catchment modeling complexities, which are often entangled with associated scaling issues between catchments as one of the Two Clouds in hydrology (Beven, 1987). Traditional hydrological models need to either be simplified or made more complex to account for scaling transformations between catchments that have different complexities. This can be done by reducing or increasing the number of hydrologic parameters, which can also be reasonably interpreted as the dimensionality of static vectors. A recognized optimal d in deep learning models illustrates that the level of an appropriate scale for regionalization exists and the involved cross-catchment hydrologic complexities exceed what the physical descriptors can provide.

The performance of the CT-LSTM with one-hot vectors (CO) is slightly better than the CT-LSTM with Gaussian vectors (CG-16) as shown in Table 3, 7. To the EA-LSTM, Gaussian vector is a more suitable choice yielding better performance. Although the specific random vector strategy is favored by different LSTM choices, the outperformed random vector strategy all enjoys an advantage of high dimension characterization. On a continental scale for 531 catchments, the one-hot vector is a vector with a length of 531, while the optimal-d is 512, both of which far exceed the 27-d physical descriptors. The high-dimensional static vector \mathbf{x}^s enhance the LSTM's ability to learn across basins to a similar or even better extent than what the 27-d catchment physical descriptors are capable of performing. This insight and discovery has a broad and significant implication for hydrologists to examine the value of \mathbf{x}^s (either physical descriptors or random vectors) that were brought in for modeling catchment complexities. Specifically, a more relevant hydrologic question to ask is: "Are catchment physical descriptors sufficient to model streamflow generation complexities and gauged catchment systems? If not, how many dimensions do we need?"

5.2 High Dimensional Catchment Characterization for Regionalization

The performance using 27-d physical descriptors is slightly worse than that of the 512-d random vectors. We interpret this to mean that the high dimensional characterization of catchments benefits regionalization performance. This idea is further supported by the result in Section 4.5, where we see a certain subset of 27-d physical descriptors is also outperformed by 512-d Gaussian vectors. When physical descriptors are incomplete, catchments are less distinguishable from each other and the regionalization performance worsens, as shown in Figure 14. Also note that Kratzert et al. (2019a) pointed out that the 27-d physical features utilized in their study are intrinsically uncertain since spatial heterogeneities are simplified as spatial averages and therefore lose certain regional

information. Uncertainties in physical descriptors might be another source that downgrades the distinctiveness across basins.

Therefore, to create a system where a system of catchments can be more distinguishable from each other, or the complexity of catchment systems can be more sufficiently quantified, we showed two strategies to expand the dimension of physical descriptors, the mixed Gaussian vector (Section 4.6.1) and the use of additional embedding (Section 4.6.2).

The mixed Gaussian vector concatenates physical descriptors with additional Gaussian vectors. The added Gaussian vectors do not have any physical meaning and only fill the expanded dimension with a Gaussian random number. Overall, with physical descriptors, this high-dimensional mixed static vector preserves the physical hydrological information and the randomness simultaneously. Results (Figure 15, Table 9) in section 4.6.1 show that the high-dimensional mixed Gaussian vectors effectively distinguish catchments and thus improve the regionalization. The peak performance is realized by 512-d, which shows the largest NSE score improvement. The results also indicate that mixed Gaussian vectors always outperform pure Gaussian vectors. Given the same dimension of static vectors, the information contained in 27-d physical features improves regional modeling. In contrast to a pure random system formed by all dimensions of Gaussian vectors, the mixed Gaussian vectors introduce ordered information and physical similarities, and thus benefit regionalization.

Inserting an embedding layer before the input gates allows an opportunity to learn more information in \mathbf{x}^s as \mathbf{x}^s is transformed into a high-dimensional space (512-d). When \mathbf{x}^s is physical descriptors (PEA), its performance is approximate to the EG-512 and also much better than the EP. After the transformation, the information of the original 27-d physical descriptors is extracted in a way that benefits regionalization. Meanwhile, this performance improvement is also recognized to the case where \mathbf{x}^s is 27-d Gaussian vectors (REA). Note that the modeling capacity for both PEA and REA is identical since both have exactly the same amount of training parameters. This observation illustrates that, within a context of the gauged basin scenario, a higher dimensional space with optimal d-dimension where catchments become more distinguishable will always elevate the regionalization performance regardless of how such a high dimension space originates (either from physical descriptor or Gaussian vector).

5.3 Implications for Catchment Systems

Besides the discussion on evaluating the effectiveness in random vectors and the high dimensional characterization advantage, the random vector approach itself and its mapping mechanisms has significant implications for understanding the current work of modeling hydrologic regionalization using the LSTM based models.

Both the Gaussian vectors and the one-hot vectors map catchments into a high dimensional space. The Gaussian vector characterizes catchments as statistically independent from each other. For the one-hot vectors, the static vectors of catchments are orthogonal to each other. Although these random vectors do not quantify catchment similarities, they ensure catchments are different from each other in a consistent way. This implies that distinguishing catchments in a gauged system achieves state-of-the-art regional modeling performance in a deep learning framework, which reflects a recently raised hydrologic concern expressed by Beven (2020) “*When essential catchment characteristics are not well understood or defined and thus not even included in catchment physical descriptors, how could a derived deep learning model perform satisfactory regionalization performance?*” Although not explicitly defining catchment characteristics, our proposed random vectors can be interpreted as high-dimensional non-physical descriptors characterizing the complexity of catchments. Catchment systems are composed of linked components representing the functional relationships between weather inputs and

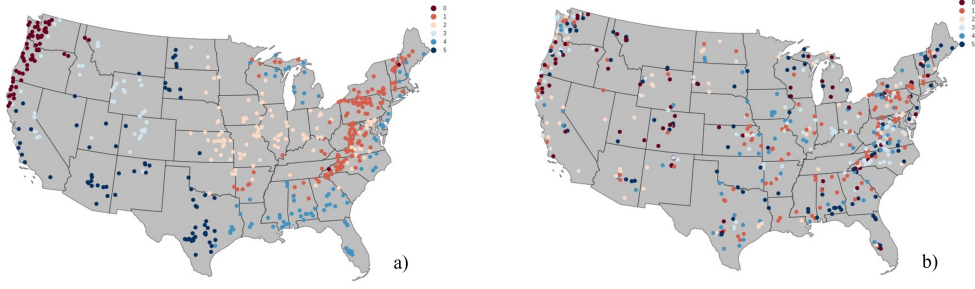


Figure 16: Clustering maps for the embedding layer of the LSTM using (a) 27-d physical descriptors and (b) 512-d Gaussian vectors. The number of cluster is six. (a) shows the clustered embedding of EP, which was also shown in the Figure 11.b in Kratzert et al. 2019a. (b) shows the clustering analysis of the EG-512 embedding.

streamflow. Arguably, a catchment system involves organized complexities where complexity may be manifested by randomness (Nearing et al., 2020; Dooge, 1986; Weinberg, 2001). The deep learning framework leverages this "random" complexity for streamflow prediction and it benefits regionalization performance to a similar extent of (if not worse than) what physical descriptors provide, which is also recognized by the observation that both PEA and REA give similar streamflow prediction performance.

Although the random vector distinguishes catchments effectively and the corresponding regionalization performance is similar to physical descriptors, it is not sufficient to self-prove the catchment uniqueness and it only serves as a surrogate of physical descriptors. Discussion on catchment uniqueness exceeds the scope of this paper. Our results imply insights to understand hydrologic similarities within a system of catchments. We analyzed and compared the patterns in the input gate (equation 9) of the EG-512 (the EA-LSTM using 512-d Gaussian vector) as well as those in the EP (the EA-LSTM using physical descriptors). This analysis intends to assess whether the original random patterns of Gaussian vectors are transformed into regional patterns internally. To remind the readers, the input gates are an embedding layer of 256-dimension as required by the LSTM model for modeling temporal complexities, which are predetermined (See the Appendix B of Kratzert et al. (2019a)). We conducted the K-means clustering analysis and created the following map below (Figure 16). The number of clusters is set to be six as suggested by Kratzert et al. (2019a).

It was previously shown that the EP automatically learns hydrological similarities and benefits the regionalization. The embeddings of the EP show a clear regional pattern (Figure 16 (a)). By contrast, for the Gaussian vector, the learned embeddings actually exhibit non-regional patterns, which implies the presence of hydrologic similarities that physical descriptors do not capture (Figure 16 (b)).

There are two interpretations for Figure 16. First, it clearly demonstrates that random vectors are not automatically transformed to physical descriptors based patterns within the LSTM cell. Second, it implies that hydrologic similarities/differences are complex patterns that go beyond what physical descriptors can explain. For two catchments geographically located far away from each other, certain hydrologic similarities in their runoff behaviors might exist and thus cluster them into similar groups. This would not be discovered by learning based on physical descriptors. Our proposed random vectors seemingly allow an opportunity for deep learning models to automatically discover such an implicit hydrologic similarity. The current 27-d physical descriptors might not be sufficient to fully represent cross-catchment hydrologic behavior similarities that are encoded in the runoff behavior. Further research is merited to investigate catchment hydrologic

similarities within a catchment system, including but not limited to the role of physical descriptors in such a context.

5.4 Implications for Deep Learning in Hydrology Regionalization

Our results are delivered in a deep learning framework. The random vector approach exhibits the strong modeling capacity of deep learning and shows a potential solution to involving complexity into a deep learning model without explicitly incorporating hydrologic processes. This approach does not add physical process understanding into the model architecture; instead, it is developed purely from a data driven perspective. We hypothesize that an appropriate dimension that accommodates catchment complexities exists and allows deep learning models to automatically distinguish cross basin similarities and therefore benefits regional modeling.

Recognizing the feasibility of using random vectors when physical descriptors are not available, another critical thought is that the current LSTM based models might not have had an appropriate architecture to fully and explicitly leverage the physical descriptors. Physical descriptors have, by their nature, physical meanings when delivered in hydrology processes. For instance, soil porosity, which is the fraction of the soil pore space, impacts the amount of water stored in the vadose zone and thus it will determine the amount of water released from soil into the discharge. This storage-discharge process shall happen after precipitation infiltrates into the subsurface, which will often be impacted by some vegetation descriptors, such as forest fraction. In short, the use of the forest fraction descriptor in process-based models comes before that of soil porosity. However, this relationship is not explicitly modeled in the current LSTM-based model since both soil porosity and forest fraction are used in an equal manner without any precedence to distinguish their hydrologic roles. The same mis-utilization of other physical descriptors is also present in particular between geomorphology descriptors and geology descriptors. It seems natural to question the validity of how the LSTM model uses \mathbf{x}^s since the use does not explicitly reflect the hydrologic roles of physical descriptors although the LSTM might implicitly learn such a relationship considering the fact that the LSTM model outperforms physical process based models (Kratzert et al., 2019a). Therefore, future research is merited to investigate and understand how the current LSTM uses \mathbf{x}^s . It might also improve the regionalization performance to adapt the LSTM model architecture by incorporating the physical meanings of the \mathbf{x}^s inputs.

The proposed random vector approach also has practical usage to assess if any given physical descriptors are complete to model catchment systems complexities. Compared to the performance with incomplete features (climate features, geology features, or geomorphology features) and to the performance without any physical descriptors, the predictive performance of the random Gaussian vectors method significantly outperforms in those scenarios. Random Gaussian vectors enable deep learning models to learn complexities more sufficiently than those physically limited descriptors. This insight has practical utility for determining the sufficiency of physical descriptors in the real world, which is challenging considering the uncertainties and complexities in hydrologic processes. When LSTM models using a specified set of physical descriptors are outperformed by random vectors, it demonstrates that those given physical descriptors are not able to resolve catchment complexities and thus suggests a need to complement them with missing features for regional modeling. For instance, as a direct illustration, Figure 4 and Table 3 suggests that 27-d physical descriptors partially address hydrologic complexities and need a certain degree of feature augmentation.

5.5 Limitations and future direction

Although the predictive performance of random vectors proves to be comparable to 27-d physical descriptors, we want to emphasize that this result is limited to gauged

prediction. The deep learning model has to have training data of the basin to predict, so the scope of this research cannot not be expanded to PUB. Therefore, recognizing this limitation, it merits future research to leverage the complexity modeling capacity found in random vectors into PUB. Note that the PEA (27-d physical descriptor with 512-d embedding) is likely a suitable option to expand into PUB since it does not involve Gaussian vectors but its applicability needs further tests.

Our ability to model catchment complexities depends on the dimension of the random vector. We show the presence of an optimal larger d , which recognizes the existence of physical processes that are not characterized, we do not provide further quantitative interpretations of the optimal d . How to utilize the observation that the optimal- d of EA-LSTM is 512 for regionalization models except for the embedding (section 4.6.2)? In future studies it will be important to identify physical processes that are not captured by physical descriptors (e.g., variable recession characteristics (Beven, 2020)) and adapt machine learning models to resolve them. Furthermore, as brought out by Beven (2020), hydrological processes are unique in different catchments. Although our results do not prove this statement of catchment uniqueness, the discovery of the regionalization effectiveness brought by random vectors could arguably have scientific implications in investigating the uniqueness of catchment. Future research is merited to investigate how to use random vectors to identify the uniqueness of catchments.

The results focus on 531 basins in the United States. Their catchment area exhibits a wide range between 4 to 1980 square kilometers, which indicates strong spatial heterogeneities across catchments. An interesting hypothesis to test is that a heterogeneous catchment prefers high dimension Gaussian vectors to account for model complexities. To test this hypothesis it will be necessary to obtain the data from catchments expressing different levels of heterogeneities. Because catchments are naturally heterogeneous, this test will require the use of synthetic data generated by physically based hydrological models. It is hypothesized that a collection of homogeneous catchment will require fewer static vectors while a collection of more complex catchment will require many static vectors. The synthetic data set will represent a system of catchments with a controlled level of heterogeneities, which will allow an opportunity to investigate how heterogeneous and homogeneous catchment systems differentiate hydrologic regionalization and modeling complexities.

Random vectors characterize a system of basins quite distinguishable in high dimensional space. The only physically distinctive information involved becomes weather inputs and associated catchment responses. This insight suggests the possibility of learning catchment similarities from weather inputs and is thus closely related to the inverse modeling problem, a field where machine learning is also advancing (Ongie et al., 2020)(Tayal et al., 2022). It therefore merits future research for an improvement in unveiling catchment characterization mysteries in a physically consistent way, likely inferred from weather inputs and catchment responses.

Our discovery has strong generalizable implications for other applications in water related or science problems. Regionalization can be conceptualized in a broader concept, that is, each local entity contributes to learn a regional or global model where cross entity information sharing benefits the predictive performance. In the context of stream-flow prediction, an entity is a catchment. For water science, an entity can also be a reservoir, lake, stream, etc. The target variable might vary depending on specific problems to solve where each problem may require a different set of entity descriptors. Mathematically, entities can be approximate functions in identical formulations with varying parameters. The benefit of random vectors in modeling regional complexities merits further research to demonstrate their practical applicability. We expect further research can test our proposed random vector approach to solve general regionalization problems across disciplines.

6 Conclusion

In this work we showed that random vectors can be used for hydrologic regionalization when catchment physical descriptors are not available. Random vector based hydrologic regionalization shows robust performance even under data sparsity and different model strategies. This method can also identify if any given physical descriptors are sufficient to account for rainfall runoff complexities. In summary, the scientific contributions of this paper are:

- The random vector method was proposed and used for regionalization in the absence of explicit physical descriptors.
- Random vectors show robust performance even under different data sparsity scenarios and different LSTM based model selection.
- Characterizing catchments in high-dimensional characteristics will improve regionalization performance.
- Random vectors can improve streamflow prediction when insufficient and uncertain basin characteristics are hard to distinguish basins. Thus, random vectors have a practical usage in determining if any given physical features are sufficient.

We also investigated scientific implications of the dimension of random vectors. This provides useful insights for the development of hydrologic models to address the model complexity and associated scaling issues.

Acknowledgement

The authors declare that they have no conflict of interest. This work was funded by the NSF HDR Grant: NSF Award 1934721. J.L. Nieber’s effort on this project was partially supported by the USDA National Institute of Food and Agriculture, Hatch/Multistate project MN 12- 109. Access to computing facilities was provided by the Minnesota Supercomputing Institute (<https://www.msi.umn.edu/>). This paper relies on open-source software and all programming was done in Python environment (Van Rossum & Drake, 2009) and its relevant packages, such as pytorch (Paszke et al., 2019),

Data Availability Statement

Both the CAMELS data (<https://doi.org/10.5065/D6G73C3Q>) and the extended Maurer forcing data (<https://doi.org/10.4211/hs.17c896843cf940339c3c3496d0c1c077>) are publically available. The code to reproduce our work and results is available at (doi.org/10.5281/zenodo.6960022).

Author Contribution

XL and AK had the idea for Gaussian vectors. VK had the idea for one-hot vectors. All the authors were involved in the discussion of experiments design and results, which were mainly led by XL and AK. XL conducted all the experiments and analyzed the results together with AK. KT supervised the experiment in Section 4.6.2. XL, AK, XJ, KC, JN, CD, MS, VK worked on the manuscripts. XL wrote the original draft and led the editing. JN, CD supervised the manuscript from the hydrologist perspective. AK, KC, XJ, MS, VK supervised the manuscript from the computer scientist perspective.

Appendix A physical descriptor description (CAMELS)

Table A1: 27-d physical descriptors in CAMELS. Descriptions are from (Addor et al., 2017)

Category	Physical descriptors	Description
climate (9)	p_mean	Mean daily precipitation
	pet_mean	Mean daily potential evapotranspiration.
	aridity	Ratio of mean PET to mean precipitation.
	p_seasonality	Seasonality and timing of precipitation. Estimated by representing annual precipitation and temperature as sine waves. Positive (negative) values indicate precipitation peaks during the summer (winter). Values of approx. 0 indicate uniform precipitation throughout the year.
	frac_snow_daily	Fraction of precipitation falling on days with temperatures below 0 .
	high_prec_freq	Frequency of high-precipitation days (≥ 5 times mean daily precipitation).
	high_prec_dur	Average duration of high-precipitation events (number of consecutive days with ≥ 5 times mean daily precipitation).
	low_prec_freq	Frequency of dry days ($< 1 \text{ mm } d^{-1}$).
	low_prec_dur	Average duration of dry periods (number of consecutive days with precipitation $< 1 \text{ mm } d^{-1}$).
Geomorphology(8)	elev_mean	Catchment mean elevation.
	slope_mean	Catchment mean slope.
	area_gages2	Catchment area.
	forest_frac	Forest fraction.
	lai_max	Maximum monthly mean of leaf area index.
	lai_diff	Difference between the max. and min. mean of the leaf area index.
	gvf_max	Maximum monthly mean of green vegetation fraction.
	gvf_diff	Difference between the maximum and minimum monthly mean of the green vegetation fraction.
Geology(10)	soil_depth_pelletier	Depth to bedrock (maximum 50 m).
	soil_depth_statsgo	Soil depth (maximum 1.5 m).
	soil_porosity	Volumetric porosity.
	soil_conductivity	Saturated hydraulic conductivity.
	max_water_content	Maximum water content of the soil.
	sand_frac	Fraction of sand in the soil.
	silt_frac	Fraction of silt in the soil.
	clay_frac	Fraction of clay in the soil.
	carb_rocks_frac	Fraction of the catchment area characterized as “Carbonate sedimentary rocks”.
	geol_permeability	Surface permeability (log10).

Appendix B FM-LSTM

FM-LSTM uses the feature modulation concept as another modelling approach. The key idea here is to use a separate gate that takes static features as input and generates a modulation vector to modulate (adapt) the features learned by a traditional LSTM. By contrast, the FM-LSTM performs the weakest modulation since the \mathbf{x}^s is only involved for updating the hidden representation, which is the last step in an LSTM update cycle before proceeding to next timestamp.

FM-LSTM

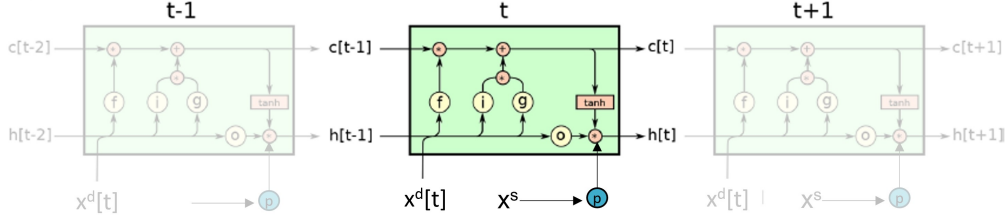


Figure B1: FMLSTM illustration, which is based on the LSTM family illustration from “Towards learning universal, regional, and local hydrological behaviors via machine learning applied to large-sample datasets” by Kratzert et al. (2019a), *Hydrology and Earth System Sciences*, 23, 5092 (Kratzert et al., 2019a)

As illustrated in Figure B1, \mathbf{x}^s is mapped to an embedding layer customized for each basin (equation B6). This is then used to modulate the hidden states output (equation B7). \mathbf{x}^s does not participate in the calculation in $\mathbf{i}[t]$, $\mathbf{f}[t]$, $\mathbf{g}[t]$, $\mathbf{o}[t]$, or $\mathbf{c}[t]$.

$$\mathbf{i}[t] = \sigma(\mathbf{W}_i \mathbf{x}^d[t] + \mathbf{U}_i \mathbf{h}[t-1] + \mathbf{b}_i) \quad (\text{B1})$$

$$\mathbf{f}[t] = \sigma(\mathbf{W}_f \mathbf{x}^d[t] + \mathbf{U}_f \mathbf{h}[t-1] + \mathbf{b}_f) \quad (\text{B2})$$

$$\mathbf{g}[t] = \tanh(\mathbf{W}_g \mathbf{x}^d[t] + \mathbf{U}_g \mathbf{h}[t-1] + \mathbf{b}_g) \quad (\text{B3})$$

$$\mathbf{o}[t] = \sigma(\mathbf{W}_o \mathbf{x}^d[t] + \mathbf{U}_o \mathbf{h}[t-1] + \mathbf{b}_o) \quad (\text{B4})$$

$$\mathbf{c}[t] = \mathbf{f}[t] \odot \mathbf{c}[t-1] + \mathbf{i}[t] \odot \mathbf{g}[t] \quad (\text{B5})$$

$$\mathbf{p} = \sigma(\mathbf{W}_p \mathbf{x}^s + b_p) \quad (\text{B6})$$

$$\mathbf{h}[t] = \mathbf{p} \odot \mathbf{o}[t] \odot \tanh(\mathbf{c}[t]) \quad (\text{B7})$$

Table B1: Random vector comparison in the FM-LSTM structure

models	Mean	Median
FP	0.653	0.698
FO	0.716	0.746
FG-512	0.695	0.738

For the FM-LSTM, we specify the optimal Gaussian vector dimension as the same of the EA-LSTM because they share the similar model modulation strategy, that is, static vectors enter the LSTM separately from the dynamic weather inputs. Using 27-d physical descriptors, Figure B2a illustrates that the FP yields worse prediction performance

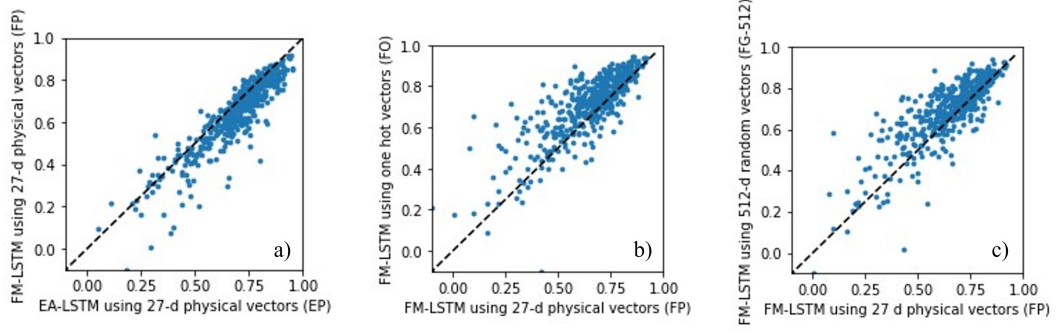


Figure B2: Predicted performance comparison of a random vector implementation in the FM-LSTM (FO and FG-512) in contrast to the FM-LSTM using 27-d physical descriptors (FP). (a) is the NSE score scatter plot of FP and EP. (b) shows the comparison between FO and FP, while (c) shows the comparison between FG-512 and FP.

compared to the EP. Even so, the FM-LSTM also attains benefits performance improvement from random vectors. Both one-hot vector and Gaussian 512-d vectors lead to significantly better predictive performance. In terms of the median, in contrast to the FP, FO elevates the performance from 0.698 to 0.746, while FG-512 improves the performance to 0.738. The one-hot vector benefits are more pronounced than those of 512-d Gaussian vectors in FM-LSTM. (Håkanson & Karlsson, 1984)

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