ML-aided Cross-Band Channel prediction in MIMO systems

- ML-stödd prediktion av kanal i tvärband i Mimo-system

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Abstract

The wireless communications technologies have experienced an exponential development during the last decades. 5G is a prominent exponent whose one of its crucial components is the Massive MIMO technology. By supporting multiple streams of signals it allows a revamped signal reconstruction in terms of mobile traffic size, data rate, latency and reliability. In this thesis work, we isolated this technology into a SIMO approach (Single-Input Multiple-Output) to explore a Machine Learning modeling to address the so-called Channel Prediction problem. Generally, the algorithms available to perform Channel Estimation in FDD and TDD deployments incur computational complexity downsides and require explicit feedback from client devices, which is typically prohibitive.

This thesis work focuses on Channel Prediction by aims of employing Machine and Deep Learning models in order to reduce the computational complexity by further relying in statistical modeling/learning. We explored the cross-Frequency Subband prediction intra-TTI (Transmission Time Interval) by means of proposing 3 three models. These intended to leverage frequency Multipath Components dependencies along TTIs. The first two ones are Probabilistic Principal Components Analysis (PPCA) and its Bayesian counterpart, Bayesian Principal Components Analysis (BPCA). Then, we implemented a Deep Learning Variational Encoder-Decoder (VED) architecture. These three models intended to deal with the hugely high-dimensional space of the 4 datasets used by its intrinsic dimensionality reduction. The PPCA method was in average five times better than the VED alternative in terms of MSE accounting for all the datasets used.

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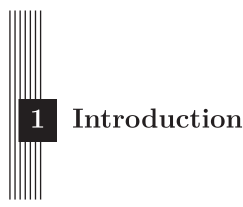
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1.1 Motivation

The exponential development in the field of wireless communications during the last decades has generated a well-established trend in the research of 5G, and WiFi technologies. To meet the 5G requirement demands in terms of performance, data-rate, spectral efficiency and network capacity, Multiple-Input Multiple-Output (MIMO) is promoted as a promising field of research to be explored. Such strategies aim at increasing the throughput and capacity of wireless networks addressing the issue of multi-antenna approaches, thus taking advantage of more data streams and signal quality [1].

A common wireless transmission context is encompassed in an imaginary portion of the land named Cellular Network (Cell). This, in turn, includes a Base Station (BS) with a geometrically-arranged number of antennas (in the order 64 to 128 for massive-MIMO). This BS communicates wirelessly with different user equipment (UE) typically in movement. The electromagnetic (EM) signal sent is affected in multiple ways when traversing the wireless environment in $BS \to UE$ (called Downlink), and $UE \to BS$ (called Uplink) directions [2].

Characterising the effect of this wireless environment in the radio signal, also known as Channel, is a core problem addressed in the field of telecommunications, the so-called Channel Estimation. In addition, when applied in multiple-antenna scenarios as such, this procedure generally leads to high overheads and computational complexity, therefore is deemed a prohibitive task in its related scientific literature. To circumvent this drawback, numerous strategies have been studied using a wide variety of approaches (see Literature Review in Section 2) among which we highlight the Channel Prediction through Statistical Modeling or Machine Learning (ML) techniques. When tackling the Channel Prediction problem, it is intended to save computational resources and overcome the limiting time constraints of numerical optimisation techniques. The R&D Center of Huawei Technologies Sweden AB is regarded as prominent in such a research domain, being selected by ICT China 2021 as the "Best Solution Case" at the PT Expo China (PTEXPO) 2021 for its outstanding network performance and extensive commercialisation of indoor distributed Massive MIMO [3]. This thesis is done in collaboration with Huawei Technologies Sweden AB due to their long-standing interest in applying ML techniques in pursue of a revamped Channel Prediction.

1.2 Background

The aforementioned wireless signal undergoes fading, that is to say, random variations in the attenuation of the signal transmitted. This is generated by the Multipath Propagation phenomenon (among others reasons like power decay), that makes the signal arrive by different separate subpaths caused by the reflection and refraction against geographical elements (scatterers) in its way. The Multipath Propagation explains shifts in phase (time delays) and amplitude (interferences) of the radio signal, specified as τ and α respectively in Figure 1.1, known as Multipath Components (MPC). This issue is of utmost importance in our thesis, because when the signal is received in BS, the different subpaths can be leveraged to reconstruct the original signal with more quality. Note that some received signal paths can actually be the resulting combination of different subpaths that arrive simultaneously. This capacity of the BS of being able to discern between more similar subpaths depends on its Spatial Resolution.

In our context, this scenario will be tackled with a Single-Input Multiple-Output (SIMO) approach, since our provided data accounts for one single UE (User Equipment) and a multiple antennae set (M antennae) that receive the signal as BS (see Section 3 for complete description of the data).

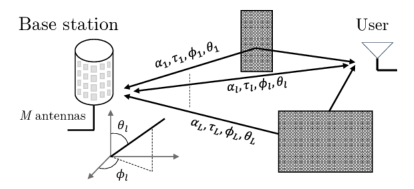


Figure 1.1: MIMO Multipath Propagation Drawing Massive MIMO multipath propagation environment described by L components. Each multipath component is characterized by its complex gain α_1 , its delay τ_l , angle to the horizon (azimuth) ϕ_l and elevation angle θ_l (obtained from [4]).

The so-called channel is theoretically formulated in the following vectorial form (in complex domain), see Equations 1.1 and 1.2 below. We can distinguish an output signal assigned to a vector Y, whose $y_i, ..., y_M$ elements correspond to the complex subpath signal received in each of the M antennas. Then, the input signal X is operated on by the channel H (whose shape is analog to Y), with the presence of Additive White Gaussian Noise (AWGN) and further interference referred to as n. This H is also referred to as Channel State Information (CSI) in the literature, and describes how the signal undergoes fading, reflection on scatterers, power decay, etc. and is used to reconstruct the signal through data processing algorithms.

$$Y = HX + n \tag{1.1}$$

$$\begin{bmatrix} y_1 \\ \vdots \\ y_M \end{bmatrix} = \begin{bmatrix} h_1 \\ \vdots \\ h_M \end{bmatrix} x + \begin{bmatrix} n_1 \\ \vdots \\ n_M \end{bmatrix}$$
 (1.2)

Reconstructing the input signal by performing

$$X = H^{-1}Y \tag{1.3}$$

is considered an intractable problem to target because of the shift in H when exiting the coherence time and the computational unfeasibility of inverting it. Hence, what is done in practice is to estimate this channel characteristics via numerical/statistical procedures. This is done by periodically estimating the channel with a signal easier to decode (X_{pilot}) . This is to say, X_{pilot} is sent out and is compared with the received Y_{pilot} , then we characterise the CSI . In this context, X_{pilot} could appear in the form of Channel State Information Reference Signal (CSI-RS). This is CSI-RS is a type of pilot signal that BS sends to UE, allowing the UE to compute the Channel State Information (CSI) and deliver it back to the BS. This process allows the network to have their connected mobile devices to be coordinated [6], because with the CSI estimated, beamforming techniques can be applied for a better addressing of the signal as well as better decoding of X.

In our scenario, the channel prediction problem addresses the task of trying to get better estimates of this H in the present time (in other Frequency Subbands, with cross-frequency prediction) and in the future (for all the Subbands, with cross-time prediction). This would allow for a better beamforming in the Downlink transmission when sending the signal from the BS to the UEs in posterior stages. As shown in Figure 1.2, in our case, data arrive to BS in a pattern-based fashion. For each Transmission Time Interval (TTI), a CSI matrix H^{SB_i} is received in one out of four Frequency subbands (SBs). We call the Frequency subband that receives H^{SB_i} pilot-receiving subband. By means of undisclosed compressing algorithms, a set of Multipath components MPC^{SB_i} are obtained by processing H^{SB_i} . That means: for each of the blue pilot-receiving subbands SB_i in each TTI, we are provided with a set of MPC^{SB_i} of size L (see Figure 1.3).

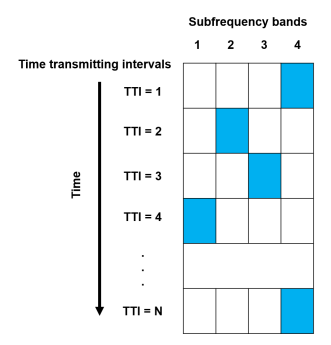


Figure 1.2: Shape of the data in a real transmission scenario (own elaboration) The blue tiles represent the pilot-receiving Frequency Subbands whereas the white tiles correspond to the pilot-missing subbands (for each TTI). Each row of data (TTI) contains $L \times 4$ Multipath Components, belonging to the four different SBs.

¹Period of time that can be calculated where the channel is considered to vary smoothly due to the absence of quick and drastic changes in the physical environment [5].

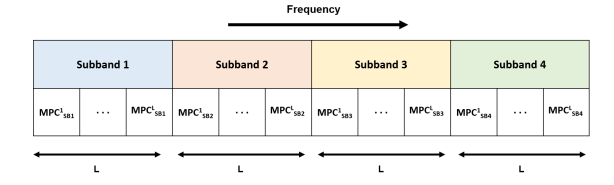


Figure 1.3: Shape of one sample TTI of the data (own elaboration).

1.3 Aim

The main objective of this thesis work is to train a model that serves us to predict the MPC^{SB_i} outside the pilot-receiving subband for each time sample TTI. This is to be performed by leveraging the dependencies in the frequency domain (across frequency subband) and/or time domain samples (across TTI). Predicting the three missing MPC^{SB_i} for each TTI leads to the reconstruction of the signal, the analog missing H^{SB_i} . We want to rely on Machine Learning and Statistical Modeling of the data in order not to incur complex and slow channel estimation algorithms. Building upon this core objective, we derive other sub-objectives:

- Characterisation of a predictive distribution $p(x_{missing} \mid x_{observed}, \theta)$, where $x_{observed}$ are the observed MPC_i^{SB} we train the model with (in pilot-receiving subbands), whereas $x_{missing}$ are the MPC^{SB_i} we want to predict (in pilot-missing subbands). Finally, θ are the trained parameters of the ML-model. In overall, performing a cross-band prediction intra-TTI. Some sub-objectives arise from this main one:
 - Visually inspect the predictions to evaluate if the distribution of the reconstructed data has been captured properly.
 - Reconstruct the TTI samples leveraging also the dependencies across time (inter-TTI).
 - Evaluate the prediction with several error metrics:
 - * Assess the quality of the prediction with statistical-based error metrics: MSE (Mean Squared Error), RMSE (Root Mean Squared Error) and MAE (Mean Absolute Error).
 - * As the predicted CSI is used in this uplink stage for the further improvement of downlink transmission, usage of downlink reconstruction transmission metrics to assess the quality of the prediction (with a reconstructed *H* matrix). Some recurrent metrics of this kind are: Beamforming gain, Network capacity and SNR (Signal-to-Noise ratio) gain.

1.4 Delimitations

One possible delimitation is the amount of data at our disposal. We have to deal with such a high dimensionality feature-wise, having in some datasets 100 times more features than datapoints. For this reason we consider the main drawback is the data quantity, as it would be a more ideal scenario to have 10 times more datapoints than features.

Moreover, the UE simulated data we have to deal with come from very stochastic simulated processes that are difficult to map to straight forward statistical distributions. That prevents us from having a complete understanding of how the data is distributed over the features and time domain beforehand. Every simulated UE is unique and we assume it is unlikely to being able to train a model that can generalize the prediction for more than one intended UE (multi-UE model prediction). Therefore, we modeled each UE per separate (with one merged dataset as an exception, see Section 3 for the data description). However, we suspect a study with larger amount of data could achieve to train a model with a training dataset consisting of different, mixed-up UEs and predict more accurately for different UEs in the testing sets.

From a benchmarking perspective, another limitation of the work is that our results are not fully comparable to the related work described in Section 2. This is because we use metrics like MSE and MAE while the gold standard metrics used in the related literature are channel gain, beamforming gain, SNR gain, etc. We deal MPC data while the mentioned works use the channel matrix data for each TTI. In order to obtain these wireless engineering metrics it would be necessary to reconstruct the channel matrix in each TTI by applying further signal processing on the MPCs. Furthermore, in order to obtain wireless engineering metrics like beamforming gain it is necessary to run many experiments by simulating more data at different Signal-to-Noise-Ratios, unfeasible for the scope of this thesis.



Literature Review

2.1 Related Work

During this section we firstly present a thorough literature review to describe how the related research works have tackled this issue of channel prediction. Various authors have been working in this matter using different approaches and techniques.

2.1.1 Signal processing approach

There exist a non-ML classical approach that tries to predict future channel CSI relying on signal processing techniques based on numerical/statistical optimization. Before addressing the cutting-edge signal processing techniques for channel prediction we should first mention some classical signal processing works. There is an example that can showcase a comprehensive benchmark for an easier but intrinsically related task, the Channel Estimation (CE). In this regard, Ma et al. [7] used the Discrete Fourier Transformation (DFT) to estimate the noise variance of the channel and a Channel Autocorrelation Matrix. Then, the Minimum Mean Squared Error (MMSE) technique is used for the Orthogonal Frequency Division Multiplexing (OFDM) to obtain a proper estimate on the channel (H_{MMSE}) with aim to decode the sent information (X) as approximately as possible. Note that this mentioned work encompasses the traditional tools (eg, Fourier Transform and MMSE) to address the problem of CE whose complexity is inferior to the following Channel Prediction techniques. Remember the Fourier Transform maps a function of time (x(t)), to a function of frequency $(X(\omega))$.

In matters of channel prediction, Swarun [8] introduced R2-F2, a system that enables LTE base stations to infer the downlink channels to a UE by observing the uplink channels from it through optimisation techniques. It is regarded as the State-of-the-Art (SOTA) when it comes to channel prediction by derivation from uplink to downlink without feedback intended in FDD MIMO systems. Wong & Evans [9], drew on non-iterative eigen-analysis technique Estimation of Signal Parameters via Rotational Invariance Techniques (ESPRIT).

Peng et al. [10] used a first-order Taylor expansion-based channel prediction model to characterise the channel. They also proposed a prediction scheme that features estimation and prediction sections and thus deriving interval of effective predictions (IEP). They evaluate the model using MSE varying SNR.

In general, techniques such as Multiple Signal Classification (MUSIC) and Estimation of Signal Parameters by Rotational Invariance Techniques (ESPRIT) lack of generality and applicability as they are sensitive to the change of the data and hardware characteristics. Their estimation process is generally tedious and computationally complex due to the manipulation of high-dimensional matrices. Finally, their obtained estimates are short-lasting due to variant nature of propagation environments with moving UEs [11].

2.1.2 ML approach

2.1.3 Cross-Temporal prediction

The attempt of channel prediction across time is another among the most wide-spread approach in the literature review collected. We can distinguish the scientific papers by the nature of the models used. Regarding time series model approach, we should point out the auto-regressive (AR) and Kalman Filtering-related scientific works, sometimes in combination with DL or ML. The mentioned techniques in this subsection are regarded as the SOTA in this research track. Jiang & Schotten [12] aim at predicting future CSI from its past values at the same frequency/sub-carrier. They used Kalman Filters to estimate AR coefficients and also a RNN-based predictor. They evaluated the performance on computational complexity with a SNR gain of 5dB at $Pr(R) = 10^{-3}$, also with an error of MSE = -44.8dB.

Liu et al. [11] made usage of Fully Connected Recurrent Neural Networks (FCRNNs) for narrow-band channel prediction. They used three different algorithms: the Real Time Recurrent Learning (RTRL), the Global Extended Kalman filter (GEKF) and the decoupled extended Kalman filter (DEKF) in order to train the Recurrent Neural Network (RNN) based channel predictor. GEKF and DEKF proved to converge faster than RTRL.

Kim et al. [13] developed and compared a vector Kalman Filter (VKF)-based channel predictor and a Machine Learning (ML)-based channel predictor using the realistic channels from the spatial channel model (SCM). The VKF-based channel predictor exploited the Auto-Regressive (AR) parameters estimated from SCM channels based on the Yule-Walker equations. Then, a MLP-based predictor was trained with the already processed data. Performance of both approaches was assessed through NMSE over different SNR. VKF's complexity was generally smaller, however when the MLP model was trained its complexity was inferior.

There also exist a well established approach where the reliance of time series forecasting on Neural Networks is stronger. In this sense, Lemayian & Hamamreh [14] proposed a Recurrent Neural Network (RNN) with Long-Short Term Memory cells (LSTM). They aimed at demonstrating low complexity channel predictions. With a similar approach but with more complex architectures, Wang et al. [15] proposed a Convolutional Long Short Term Memory Network (ConvLSTM-net)-based method. Featuring a 5x5 patch, this architecture achieved around NMSE = -37 dB in both time and frequency domain. In this line, Yuan et al. [16] combined pattern extraction performed by a convolutional neural network (CNN) and a CSI predictor configured as an auto-regressive (AR) model. They also used an ARNN (Auto-regressive Neural network) as an alternative of the AR-model as CSI predictor. In the first stage the pattern was identified and the latter step yields the prediction. They evaluated models' performances through NMSE over P (order of AR parameters).

Some other works addressed this problem deviating from the AR modeling and focusing in dealing with the interactions of complex variables. For instance, Ding & Hirose [17] presented a method for predicting time-varying channels by combining a multilayer complex-valued neural network (CVNN) with the chirp z-transform (CZT), resulting in a CZT-ML-CVNN-based predictor. They present the accuracy in terms of SNR vs Bit Error Rate, where CZT-ML-CVNN-based outperformed the other architectures. Similarly, Yang et al.[18] used sparse complex-valued neural network (SCNet) to leverage the uplink CSI to predict the downlink

feedback motivated by the universal approximation theorem. They relied on NMSE metric over Δf and AS (Angle Spread).

2.1.4 Cross-Frequency Band prediction

We should recall that our data is based in Multipath Components that comes out of data compression stage from CSI channel matrices. Identifying research works that exactly map our scenario and apply cross frequency subband channel prediction have not been entirely possible. In spite of that, we could collect some projects that use channel prediction across Frequency Band in Frequency Division Multiplexing (FDD) communication scheme.

Bakshi et al. [19] trained a Neural Network on a standard channel model to generate coarse estimates for the underlying variables of the channel. The NN was embedded in a system they called OptML. Their goal was to reduce the complexity of the model in an order of magnitude 10x, while maintaining similar prediction quality. A Neural Network Distance Estimator (NNDE) yielded coarse estimates for distances between MPCs, these underwent further optimization and prediction processes.

2.1.5 Our intuition on the Channel Prediction problem

Not related to the SIMO or MIMO domain, but still framed in the wireless communications realm, Frohle et al. elaborated on Gaussian Processes (GPs) in two different studies [20], [21]. They aimed at predicting the wireless channel between mobile agents incorporating the agents' location uncertainty. We would like to claim that the spatial-temporal dependencies exploited by GPs could fare well to our problem because time and frequency dependencies could be exploited. Trying to estimate such a high-dimensional kernel could perhaps be computationally problematic though.

Therefore, we implemented PPCA as a baseline model for prediction due to its low-complexity covariance estimation that entails feature selection as well. Previous works like Porta [22], used PPCA for action selection in an Active Appearance -Based Robot Location framework. They concretely imputate missing values with PPCA in order to apply this modeling to depth maps for the robot to robustly be space-aware. Nyamundanda [23] used PPCA for metabolic studies where the missing value imputation was necessary by modeling the estimated covariance between the high-dimensional metabolite features. The Bayesian PPCA (BPCA) was used in Oba et al. [24] for missing value imputation in gene expression profiling by the modeling of its latent feature space. It outperformed singular value decomposition and K -nearest neighbors for this same task. Jenelius & Koutsopoulos used a similar missing value inference in the Urban Network Travel Time Prediction Based task [25]. They modeled the PPCA through EM-algorithm in order to predict time delays of feature observations conditioned on the already observed information using the Gaussian conditional property.

A similar modeling was intended in our baseline method, this is to say, conditioning the unobserved (missing) MPCs corresponding to the pilot-missing frequency subbands on the pilot-receiving ones. Therefore, our PPCA model is to be depicted in Section 4.1.

Aiming to use a more powerful non-linear modeling, we wanted to follow the approaches of Linfu et al. [26] and Hussein et al. [27]. These works incorporated Variational Autoencoders (VAEs) to address the downlink CSI obtention problem in a frequency division duplexing (FDD)-MIMO setting. The uplink CSI can be obtained by means of Channel Estimation while the downlink CSI should be sent to the BS from the UE, which incurs a huge overhead. These works intend to solve the problem addressed to learn an optimal/accurate compression of this downlink CSI.

Likewise, we implemented a Variational Encoder-Decoder (VED) architecture pursuing to learn an optimal reconstruction mapping as described previously for the PPCA baseline model. In short, we aimed to estimate a proper latent feature space in order to reconstruct the missing MPCs we receive at the BS was meant (note these MPCs come from prior compressing stages). Thus, for this reconstruction, we need to leverage the observed information of the pilot-receiving frequency band to later extrapolate to the missing ones. This model is described in Section 4.4. We based our model implementation in Liu et al. [28], where they used a similar modification of a VAE to predict the downlink CSI from the uplink CSI in a FDD system. Note what we used data of another nature because we dealt with MPCs resulting from further signal processing of the uplink's CSI. They achieved a SNR improvement over 10 dB in MIMO transmissions compared to the current state-of-the-art in this regard, the R2-F2 technique [8].



This section outlines the data handled in our thesis work. In the Section 3.1 the data obtained is described and in Section 3.2 the pre-processing carried out is explained. In addition, we explain how further hold-out splitting and data masking is performed for the following experiments, see Section 3.4.

3.1 Data obtention

The data used in this thesis consist of three different datasets. All of them where produced by simulations carried out by the Huawei Technologies Sweden AB 's Wireless Algorithm Lab. All three datasets are three-dimensional tensors [L x SB x TTI]. The parameter TTI corresponds to Transmission Time Intervals. They are called intervals because these are the duration of the transmission which is related to the encapsulation of data in blocks for an efficient communication. In practice we can treat this dimension as simple time stamps where we receive the data in a time series fashion. SB stands for Frequency Subbands, these are always fixed to 4. L, in turn, stands for the number of Multipath Components per band, which are complex signal values.

The effective datasets used in the model executions are the ones specified in Table 3.1. We used a dataset called UE0 (90 TTIs), and two other datasets named UE1 and UE2, with 590 TTIs each. All the last 3 mentioned correspond to 3 fictional User Equipments. However, the forth dataset named JointDS is the result of the merge of UE1 and UE2. The details the dataset arrangements are specified in Section 3.2 and 3.4.

Table 3.1: Description of datasets.

Dataset Transmisison Time intervals		Multipath Components / SB	Subbands
	TTI	MPCs / SB	$_{ m SB}$
UE0	90	309	4
UE1	590	640	4
UE2	590	640	4
JointDS	1180	640	4

3.2 Data pre-processing

In order to pre-process the data for it to be fitted to a learnable ML model, we created some scripts in R Language to unfold these 3-dimensional tensors into 2-dimensional matrices. We concatenated each [TTI x L] slice of the tensors we were provided (horizontally) so as we got resulting matrices of 2 dimensions [TTI x L*4]. As every value in the dataset was in the Cartesian complex plane (z=a+bi), we separated the real part $(\Re \mathfrak{e}(z)=a)$ from the imaginary one $(\Im \mathfrak{m}(z)=b$ without the i to allow the computation) in different columns. Therefore, the final matrices are of [TTI x L*4*2]. The last mentioned process (separating $\Re \mathfrak{e}(z)$ from $\Im \mathfrak{m}(z)$ in different columns) is to allow for the implemented models to be compatible with the computation. In order to train the models, we split the matrix row-wise in a 60% / 40% fashion (Training Set/ Validation Set).

Dataset Transmisison Time intervals		Features	
	TTI	4 SB Concatenated Horizontally and Unfolded	
UE0	90	2472	
UE1	590	5120	
UE2	590	5120	
JointDS	1180	5120	

Table 3.2: Description of unfolding of datasets.

3.3 Data description

A screenshot of dataset UE0 is provided in Figure 3.1 to visually show how the data looks like. The data consists on complex amplitudes (MPCs) but with imaginary and real part separate by columns. Every row corresponds to a time sample for all the Multipath Components while every column represents one Multipath Component for a concrete band (either the real or imaginary part). When we say "features" we refer of the columns of this datasets, i.e. the real or imaginary part of a Multipath Component that evolves over time. Note that under the bracket of, for instance, Subband 1, the first column of the real part has its counterpart in the first column of the imaginary part, both conforming the first Multipath Component in the SB 1. Then, following this logic, the first column of SB2, SB3 and SB4 is the 1st Multipath Component (real part) but in the other bands.

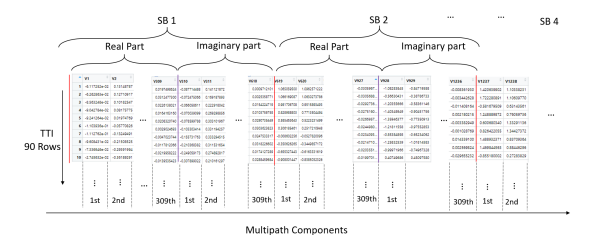


Figure 3.1: Screenshoot of the data matrix. Dataset UE0 (own elaboration).

In Figures 3.2 and 3.3 how the 1st MPC in the 4 Subbands evolves across the TTI dimension (time) in its real and imaginary part respectively. In this plots, each line represents a time series plot of a column in the dataset. Notice how the MPCs evolve in a sinusoidal pattern.

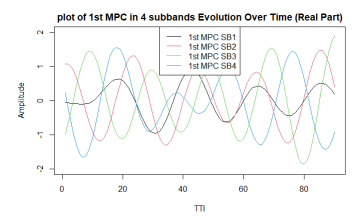


Figure 3.2: Real Part of the 1st MPC in the 4 SBs. Dataset UE0 (own elaboration).

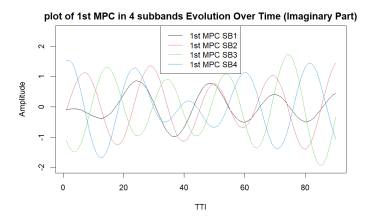


Figure 3.3: Imaginary Part of the 1st MPC in the 4 SBs. Dataset UE0 (own elaboration).

Finally, in Figure 3.4 we display the plot of one TTI, in other words, one row of the dataset UE0, (first row of the matrix in Figure 3.1).

3.4 Masking and hold-out split

As described in Section 1.2, in a real transmission situation, every TTI of data is sensed in such a way we can only observe one out of four Frequency Subbands. This masking occurs in a well-defined fixed pattern, which in this work we decided to set as " $1 \rightarrow 3 \rightarrow 2 \rightarrow 4$ " (these are the observed subbands at each TTI). For this reason, by scripting (in R), we applied the masking in 2 different ways depending on what model we made use of. When we say we mask a certain TTI (row in the dataset), we refer to fill with NaNs (Not a Number numeric data type) all of the MPCs of a certain Frequency Subband out of the four.

For both of the two models, the dataset hold-out splitting was applied respecting the time sequence, leaving the last 40% of TTIs for the validation set. See Table 3.3 for description of the hold-out split result.

For the training and validation set of the JointDS, we just merged the respective training and validation sets of UE1 and UE2, hence respecting the time-line stream in which the samples

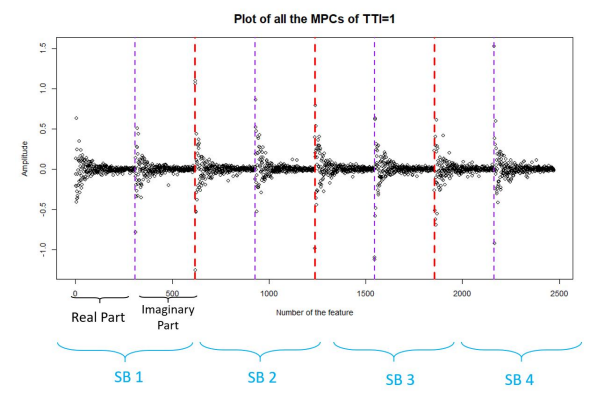


Figure 3.4: Plot of all the MPCs over one TTI, notice Real and Imaginary parts for all four Subbands. Dataset UE0 (own elaboration).

Table 3.3: Description of hold-out split.

Dataset	Training	Validation	
	TTIs	TTIs	
UE0	54	36	
UE1	354	236	
UE2	354	236	
JointDS	708	472	

where produced in a real-life situation (not shuffling TTIs between training and validation splits for none of UE1 and UE2).

3.4.1 Masking for PPCA model

None of the training datasets are masked when it comes to training the PPCA model. This decision in meant to assure that we are extracting proper W loadings that encapsulate the greatest variance in the latent space well enough. After that, we masked the validation sets in the pattern described in the previous point in order to evaluate the predictive performance leveraged from the PPCA's obtained W factor loadings.

3.4.2 Hold-out for Variational Encoder-Decoder model

For our implemented Variational Encoder-Decoder model, we masked both training and validation sets. This whole dataset masking is intended to force a supervised learning, creating (x_i, y_i) tuples that were fed to the DL architecture. For each TTI, x_i corresponds to the Multipath Components in the observed Frequency Subband, (25% of the TTI), treated as

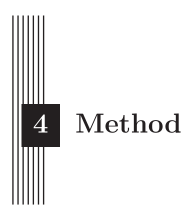
predictors. The target features, collected in y_i , correspond to the remaining 75% of the TTI, and they are the unobserved MPCs corresponding to the three unobserved Frequency SBs. These (x_i, y_i) are equivalent to the notation (X_{obs}, X_{mis}) that will be used in the Section 4.1 on PPCA theory.

The training set for every dataset is normalised to be fed into the VED model. That means subtracting the mean and dividing by the standard deviation (STD) feature-wise. We applied normalization in the validation set according to the mean and STD of the training set in order to prevent data leakage.

The Table 3.4 summarizes the masking and hold-out properties of each subset.

Table 3.4: Description of hold-out split datasets properties.

Model	Dataset	Obs(X) / Mis(Y)	Normalization	Masking
PPCA	Training		No	No
TTOA	Validation		INO	Yes
	Training	Observed	Yes	
VED		Missing	No	Yes
V ED	Validation	Observed	Yes	res
	Vandation	Missing	No	



In this section, we present the theoretical background of the models used in the thesis project. In the Sections 4.1 and 4.4, PPCA and VED are described. Note that the first subsections of the Section 4.4 constitute a preliminary Deep Learning framework explanation to make the background considerations of VAEs/VED more feasible for the reader. In Section 5, we describe how the experiments of PPCAs and VED were carried out, as well as the configured experimental set-up.

4.1 Probabilistic Principal Component Analysis (PPCA)

This point is to outline the two baseline models used in the thesis (PPCA and BPCA). In order to describe these models, we should first outline the Factor Analysis method for an intuitive build-up on this matter.

4.1.1 Factor Analysis (FA)

Let us assume a Gaussian vector x_i of dimension $S \times 1$, assume $x_i \sim \mathcal{N}(\mu, \Sigma)$. We can consider S to be the size of the total MPCs accounting for all bands (total features per row). Now let

$$x_i = \mu + W z_i^T + \varepsilon_i, \tag{4.1}$$

where μ is the multivariate mean of x_i $[S \times 1]$, ε is the error term of dimension $[S \times 1]$ and $\varepsilon \sim \mathcal{N}(0, \Psi)$, and Ψ is the variance of the noise. z_i and W have dimensions $[Q \times 1]$ and $[S \times Q]$ respectively, where Q is a parameter of design choice (number of factors). W is also known as the loading vector of all the features S (MPCs) for all the factors. Meanwhile, z_i is the score value of each of the features of x_i for each factor.

Conditioning the observations on the rest of the parameters, x_i can be modeled as

$$x_i \mid z_i \sim \mathcal{N}(\mu + Wz_i, \Psi),$$
 (4.2)

where we assume

$$z_i \sim \mathcal{N}(0, I). \tag{4.3}$$

Then, integrating the latent variable z out we yield

$$x_i \sim \mathcal{N}(\mu, WW^T + \Psi), \tag{4.4}$$

where $WW^T + \Psi = \Sigma$, the estimated covariance.

The general matrix for the entire dataset reads as follows

$$X = W Z + \mu + \varepsilon, \tag{4.5}$$

where W is $[N \times Q]$, Z is $[Q \times S]$, ε is $[N \times S]$, μ is $[N \times S]$ and it results in the reconstructed dataset X with dimensions $[N \times S]$.

4.2 Theoretical transition from FA to PPCA

In order to transition to PPCA, we add this constraint to the Factor Analysis model to obtain the PPCA model

$$\Psi = \sigma^2 I. \tag{4.6}$$

PPCA assumes that the data points are independent of each other given the latent variables and adds an orthogonality assumption in the factor loadings matrix W.

$$x_i \mid z_i \sim \mathcal{N}(\mu + W z_i^T, \quad \sigma^2 I)$$
 (4.7)

Analogue to Factor Analysis modeling, let us integrate over z:

$$x_i \sim \mathcal{N}(\mu, WW^T + \sigma^2 I), \tag{4.8}$$

where $WW^T + \sigma^2 I = \Sigma$, the estimated covariance.

We assume prior $z_i \sim N(\mu_o, \Sigma_o)$ and to simplify we can promote $z_i \sim N(0, I)$. Once we have estimated μ and W from the PPCA method we are interested in characterising the predictive distribution. Note first that the distribution of the latent variables given the observed data can be derived using Bayes' Theorem to give:

$$z_i \mid x_i \sim \mathcal{N}((WW^T + \sigma^2 I)^{-1}W^T(x_i - \mu), \quad \sigma^2(WW^T + \sigma^2 I)^{-1}).$$
 (4.9)

4.2.1 Dealing with missing data through inference on latent variables

Once the Probabilistic Principal Components (PPCs) are fitted, we followed the approach in Jenelius & Koutsopoulos [25] to attain a predictive distribution/function. For each sample x_i ([$S \times 1$]) we divide the Multipath Components into $x_{observed}$ (x_{obs}) and $x_{missing}$ (x_{mis}). We distinguish between 2 vectors of indices that discern between the observed features and the missing ones (namely Idx_{obs} , Idx_{mis} with dimensions [$S_{obs} \times 1$] and [$S_{mis} \times 1$]). These correspond to the frequency subband with received pilot and the ones with no observed pilot respectively. Therefore, we define $x_{obs} = x[Idx_{obs}]$ and $x_{mis} = x[Idx_{mis}]$.

$$\mathbf{X_{obs}} = \begin{bmatrix} x_{Idx_{obs}}[1] \\ \vdots \\ x_{Idx_{obs}}[K_{obs}] \end{bmatrix} \quad \mathbf{X_{mis}} = \begin{bmatrix} x_{Idx_{mis}}[1] \\ \vdots \\ x_{Idx_{mis}}[K_{mis}] \end{bmatrix}$$

$$(4.10)$$

Then, the conditional property of loading matrix W let us split it into the $[S_{obs} \times Q]$ matrix W_{obs} and the $[S_{mis} \times Q]$ matrix W_{mis} . Similarly, we apply the same to μ , splitting it into μ_{obs} and μ_{mis} (where Q is the number of PPCs extracted).

We use the conditioning properties of the Multivariate Normal Distribution to obtain our estimates $x_{mis} \mid x_{obs} \sim \mathcal{N}(\hat{x}_{mis|obs}, \Sigma_{mis|obs})$, such that

$$\hat{x}_{mis|obs} = \mu_{mis} + W_{mis}(W_{obs}^T W_{obs} + \sigma^2 I)^{-1} W_{obs}^T (x_{obs} - \mu_{obs}), \tag{4.11}$$

$$\hat{\Sigma}_{mis|obs} = \sigma^2 W_{mis} (W_{obs}^T W_{obs} + \sigma^2 I)^{-1} W_{mis}^T + \sigma^2 I.$$
(4.12)

The mean vector $\hat{\mu}_{mis}$ serves as point estimate for the missing Multipath Components, whereas $\Sigma_{mis|obs}$ describes the variability around those predictions. μ_{mis} , in turn, establishes a baseline for the prediction to be added to the product of the updated loadings $W_{mis}(W_{obs}^TW_{obs} + \sigma^2I)^{-1}W_{obs}^T$ times the difference vector between observed MPCs and MPCs' historical means $(x_{obs} - \mu_{obs})$.

Here below, we show how the conditional Gaussian property is applied to obtain the predictive function for $\hat{x}_{mis|obs}$. We consider a generic variable $X \sim \mathcal{N}(\mu, \Sigma)$ with some missing and observed features. Note sub-indexes $_{mo}$ and $_{om}$ stand for $_{missing|observed}$ and $_{observed|missing}$ respectively.

$$\boldsymbol{\mu} = \begin{bmatrix} \boldsymbol{\mu}_{mis} \\ \boldsymbol{\mu}_{obs} \end{bmatrix} \qquad (4.13) \qquad \boldsymbol{X} = \begin{bmatrix} \boldsymbol{X}_{mis} \\ \boldsymbol{X}_{obs} \end{bmatrix} \qquad (4.14) \qquad \boldsymbol{\Sigma} = \begin{bmatrix} \boldsymbol{\Sigma}_{mis} & \boldsymbol{\Sigma}_{mo} \\ \boldsymbol{\Sigma}_{mi} & \boldsymbol{\Sigma}_{obs} \end{bmatrix} \qquad (4.15)$$

It follows that we attain the following point conditional estimates for the $x_{missing}$ mean and covariance ruled by $p(x_{mis} \mid x_{obs}) \sim \mathcal{N}(x_{mis} \mid \mu_{mis|obs}, \Sigma_{mis|obs})$,

$$\hat{\mu}_{mis|obs} = \mu_{mis} + \Sigma_{mo} \Sigma_{mis}^{-1} (x_{obs} - \mu_{obs}), \tag{4.16}$$

$$\hat{\Sigma}_{mis|obs} = \Sigma_{mis} - \Sigma_{mo} \Sigma_{mis}^{-1} \Sigma_{om}. \tag{4.17}$$

Notice that this result is analogous to the one in Equations 4.11 and 4.12.

4.3 BPCA

Following the approach of Oba et al. [24], we propose Bayesian Probabilistic Principal Components (BPCA) as a extension of our baseline. BPCA promotes priors for $p(W, \mu, \sigma^2)$ over the parameters of the model. Then, effective dimensionality over the matrix W is intended to be obtained by means of the introduction of a hierarchical prior $p(W \mid \alpha)$ where $\alpha = \{\alpha_1, ... \alpha_q\}$. The maximum number of α hyperparameters is q = S - 1. α_i controls the inverse variance of the corresponding PPC W_i . The larger the α_i , the smaller the W_i , eventually switching it off and hence reducing latent space dimensionality.

Finally, we can derive the natural logarithm-posterior estimate of W, W_{MP}

$$\ln p(W \mid D) = L - \frac{1}{2} \sum_{i=1}^{d-1} \alpha_i ||w_i||^2 + const.$$
 (4.18)

According to the R library used in this thesis (pcaMethods [29]) BPCA combines an EM approach for PCA with a Bayesian model. In PCA, data dissimilar to the training set but close to the latent space may have the same reconstruction error. BPCA defines a likelihood function such that the likelihood for data dissimilar the training set is much lower, even if they are close to the latent space. The algorithm does not force orthogonality between loadings W. thus, factor loadings are not necessarily orthogonal.

4.4 Variational Encoder-Decoder (VED)

As for the next course of action, in order to infer to the latent representation values to approximate the probability distribution $p(X_{mis} \mid X_{obs}, \theta_{model})$, we proposed a DL approach that consists of an Encoder-Decoder Network with a Variational Inference layer in the bottleneck. The aim of this approach is to force each TTI sample to be embedded in a continuous value of a latent space. It is inspired by Variational Autoencoders [30]. The following sections are going to serve as a summary of the Deep Learning related theory for the Variational Encoder-Decoder description to be more accessible.

4.4.1 Deep Learning

The Deep Learning term refers to a segment of techniques under the umbrella of Machine Learning Methods that draw on the use of Artificial Neural Networks (ANN) by using feature learning [31]. We can distinguish two main categories, supervised learning (where inputs are labeled or coupled with a explicit target) or unsupervised (unlabeled input data). ANNs, in turn, were inspired by the information flow, processing and distribution in biological systems. To give a visual example of an NN, we show the Multilayer Perceptron (MLP) in Figure 4.1.

4.4.1.1 Multilayer Perceptron (MLP)

MLPs are the quintessential types of NNs, we base the ensuing explanations in Bishop's Pattern Recognition and Machine Learning [32]. MLPs aim at approximating a function f^* in the context $y = f^*(x)$ that maps a value x to y. Then, let the approximation function learned by the NN be f in $y = f(x; \theta)$, obtained by tweaking the set of parameters θ until they are optimal for the approximation. A simple MLP might be defined as a composition of different functions, for instance $f^{(1)}$, $f^{(2)}$, $f^{(3)}$ being chained as $f(x) = f^{(3)}(f^{(2)}(f^{(1)}(x)))$. Let us consider a MLP with two hidden layers. Let j = 1, ..., M be the set of hidden units, while k = 1, ..., M is the set of outputs. The input data X has dimensions i = 1, ..., D. In a two hidden layer NN, we can characterize the overall feed-forward function as follows:

$$y(x,\theta) = \sigma \left(\sum_{j=1}^{M} \theta_{kj}^{(2)} h \left(\sum_{i=1}^{D} \theta_{ji}^{(1)} x_i + \theta_{j0}^{(1)} \right) + \theta_{k0}^{(2)} \right). \tag{4.19}$$

We can consider $\theta_{ji}^{(1)}$ to be the weight from input dimension i to the hidden node j in layer (1). In the intermediate output we obtain a set of M linear combinations called activations: $a_j = \sum_{i=1}^D \theta_{ji}^{(1)} x_i + \theta_{j0}^{(1)}$. σ and h stand for activation functions. We depict graphically this function mapping in the MLP in Figure 4.1.

Activation functions are non-linear functions applied to the activations resulting from the layers. They are useful to restrict the activation values, and for the entire neural network

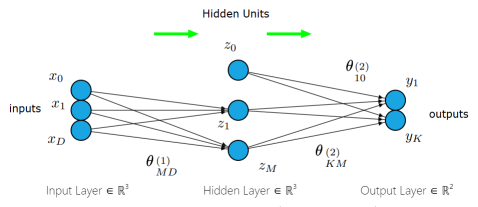


Figure 4.1: Diagram of a MLP (own elaboration).

not to collapse in a composed bigger linear function, therefore not being able to capture nonlinear patterns in the data. The following equations describe different activation functions, ReLU (Equation 4.20) sets a cut-off for negative values, hyperbolic tangent (Equation 4.21) squashes a real-valued number to the range [-1,1], softplus (Equation 4.22) can be regarded as a smooth version of ReLU and LeakyReLU (Equation 4.23) is similar to ReLU but allowing small non-negative gradients for negative values of the input.

$$ReLU(\mathbf{x}) = max(0, \mathbf{x})$$
 (4.20)

$$tanh(\mathbf{x}) = \frac{1 - e^{(-2\mathbf{x})}}{1 + e^{(-2\mathbf{x})}}$$

$$(4.21)$$

$$Softplus(x) = \frac{1}{\beta} \log(1 + \exp(\beta x)) \quad (4.22) \qquad LeakyReLU(x) = \begin{cases} x > 0 \\ \alpha x & x < 0 \end{cases} \quad (4.23)$$

4.4.1.2 Training of the Network

Neural Networks need to update their parameters so as an optimal or sub-optimal approximation is achieved by learning from a training dataset. For the sake of the explanation, we can adopt a probabilistic view of a regression problem, since the aim of the thesis is to learn a multivariate regression. Let this training dataset consist of a set of vectors $X = \{x_1...x_N\}$, being coupled with a set of target vectors $\{y_n\}$, both can take any value in real domain. We assume that y has a Gaussian distribution with an x-dependent mean, which is given by the output of the neural network, see Equation 4.24 [32]. Mind β is the inverse of the Gaussian noise (the precision). It follows that we can construct the likelihood function of the output data y_n conditioned on the input data x_n and the parameters (Equation 4.25). Finally, taking negative logarithm of the latter, we yield the error Equation 4.26.

$$p(y \mid x, \theta) = \mathcal{N}(y \mid f(x, \theta), \beta^{-1})$$
(4.24)

$$p(y \mid X, \theta, \beta) = \prod_{n=1}^{N} p(y_n \mid x_n, \theta, \beta)$$
(4.25)

$$\frac{\beta}{2} \sum_{n=1}^{N} \{ f(x_n, \theta) - y_n \}^2 - \frac{N}{2} ln\beta + \frac{N}{2} ln(2\pi)$$
 (4.26)

We should note that minimizing this expression in Equation 4.26 is the same as minimizing the squared term, which is analogue to minimizing the sum of squared errors.

4.4.1.3 Backprogation and Adam

Let then the cost function chosen for the NN's output be $E(\theta)$, dependent on the set of parameters θ . The aim is to calculate the gradient $\nabla E(\theta)$ vector that relates with the greatest increase direction. What is done in practise is decreasing in this inverse direction $-\nabla E(\theta)$ making a small step in the weight space $\theta := \theta + \nabla \theta$, where $\nabla \theta$ is the gradient vector with the corresponding magnitude to subtract to each parameter in the network. What is often carried out is gradient descent optimization, performing this update $\theta^{(\tau+1)} = \theta^{(\tau)} - \eta \nabla E(\theta^{(\tau)})$, where τ is the specific time stage and η is the learning rate (portion of the gradient to subtract). For each of the steps performed, the gradient is calculated grouping the training data into batches, thus the resulting gradient being $E(\theta) = \sum_{n=1}^{N} E_n(\theta)$. In order to update the specific weight θ , the quantity subtracted is determined by the backpropagation algorithm, described at Algorithm 1. This is to calculate the partial derivative of the weight θ_{ij} with respect to the total gradient: $\frac{\delta E}{\delta \theta_{ij}}$. Adaptive Moment Estimation (Adam) [33] is a batch-based optimisation algorithm that is computationally efficient and varies the learning rate according to the training results, see Algorithm 2.

Algorithm 1 Backpropagation Algorithm Pseudocode [34]

```
Procedure Training X \leftarrow Training Data Set of size mxn y \leftarrow Labels for records in X w \leftarrow The weights for respective layers l \leftarrow The number of layers in the neural networks, 1...L D_{ij}^{(l)} \leftarrow The error for all l,i,j t_{(ij)}^{(l)} \leftarrow 0. For all l,i,j for i=1 to m do a^l \leftarrow feedforward(x^{(i)},w) d^l \leftarrow a(L) - y(i) t_{ij}^{(l)} \leftarrow t_{ij}^{(l)} + a_j^{(l)} \cdot t_i^{l+1} end for if j \neq 0 then D_{ij}^{(l)} \leftarrow \frac{1}{m} t_{ij}^{(l)} + \lambda w_{ij}^{(l)} else D_{ij}^{(l)} \leftarrow \frac{1}{m} t_{ij}^{(l)} where \frac{\delta}{\delta w_{ij}^{(l)}} J(w) = D_{ij}^{(l)}
```

4.4.2 Probabilistic Theory of VAEs

An Autoencoder typically serves as a reconstruction-intended NN since the dimensions of the input and the output layer match and it often behaves in an unsupervised basis. Firstly, the encoder part gradually reduces the dimensionality of the input up to a bottleneck and then increases again until the input dimension is reached (both by using fully connected layers/CNN, or other sort of layers). Conversely, in our thesis, the model architecture has an input dimension of size [L] (1 Frequency Subband with its respective Multipath Components within 1 TTI)

Algorithm 2 The Adam Algorithm Pseudocode [33]

Require: Step size ϵ (Suggested default: 0.001)

Require: Exponential decay rates for moment estimates, ρ_1 and ρ_2 in [0,1). (Suggested

defaults 0.9 and 0.999 respectively)

Require: Small constant δ used for numerical stabilization. (Suggested default: 10^{-8})

Require: Initial parameters θ

Initialize 1st and 2nd moment variables s = 0, r = 0

Initialize time step t=0

while stopping criterion not met do

Sample a minibatch of m examples from the training set $\{x^{(1)},...,x^{(m)}\}$ with corresponding targets $\mathbf{y^{(i)}}$

Compute gradient: $g \leftarrow \frac{1}{m} \nabla \theta \sum_{i} L(f(x^{(i)}; \theta), y^{(i)})$

 $t \leftarrow t + 1$

Update biased first moment estimate: $s \leftarrow \rho_1 s + (1 - \rho_1)g$

Update biased second moment estimate: $r \leftarrow \rho_2 r + (1 - \rho_2) g \odot g$

Correct bias in first moment: $\hat{s} = \frac{r}{1-\rho_1^T}$ Correct bias in second moment: $\hat{r} = \frac{r}{1-\rho_2^T}$

Compute update: $\nabla \theta = -\epsilon \frac{\hat{s}}{\sqrt{\hat{r}} + \delta}$ (operations applied element-wise) Apply update: $\theta \leftarrow \theta + \nabla \theta$

and an output dimension of $[3 \times L]$ (3 remaining frequency subbands). It also has a supervised approach, aiming to reconstruct a TTI from one observed subband extrapolating to the three unobserved ones. For this reason, we have called it Variational Encoder-Decoder.

According to the formal definition in *Deep Learning* by Goodfellow [35], the encoder can be characterised as the stochastic mapping $p_{encoder}(h \mid x)$ with deterministic function h = f(x), that is the latent representation resulting in the bottleneck . The decoder, in turn, follows with $p_{decoder}(x \mid h)$ and produces a reconstruction r = g(h), that in our case is an extrapolation towards missing information in vector observation x. Please see the Figure 4.2 to visualize the main idea of the architecture.

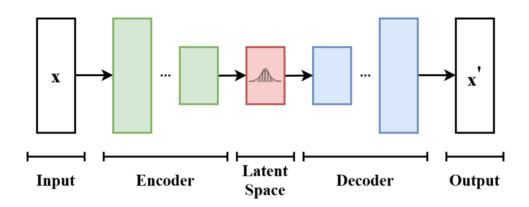


Figure 4.2: Archetypical Variational Autoencoder architecture (obtained from [36]).

Leveraging the Variational Encoder-Decoder, we aim at learning the aforementioned distributions from the training dataset. Formally, in Variational Autoencoders (according to [37]), we need to force every point of our training set X ($\vec{x}_i \in X \subset \mathbb{R}^p$) to be mapped to a vector of variables z in a lower-dimensional space \mathcal{Z} that can be sampled from P(z) defined over \mathcal{Z} . After, let us consider a family of deterministic reconstructive functions $f(z;\theta)$ parameterized by a vector θ in a space Θ , where $f: \mathcal{Z} \times \theta \to \mathcal{X}$. z is a random realization whereas θ is fixed, yielding $f(z;\theta)$ to be a random variable in the space \mathcal{X} . The model optimisation intends to tune θ so as by sampling z from p(Z), $f(z;\theta)$ can reconstruct an approximation of \mathcal{X} . In mathematical notation, let $f(z;\theta)$ be replaced by the generative distribution $P(X \mid z;\theta)$. Then, according the law of total probability, the marginal likelihood of the data is to be maximised for each X through the generative process:

$$P(X) = \int P(X \mid z; \theta) P(z) dz. \tag{4.27}$$

The typical output distribution for a VAE is a Gaussian distribution such as $P(X \mid z; \theta) = \mathcal{N}(X \mid f(z; \theta), \sigma^2 I)$, with hyper-parameter σ^2 . This decision is to increase P(X) via some optimisation technique for some sampled z, making the data more likely under the generative model.

In our scenario, we used this intuition but not in a data reconstruction manner. Indeed, we coerced the training into a supervised basis, where the data was arranged by $X_{observed}$ and $Y_{missing}$ tuples, both for the training and validation datasets. As of now, we introduce terms X_{obs} ($X_{observed}$) and Y_{mis} ($Y_{missing}$). That makes explicit that the missing MPCs for each TTI corresponding to the missing frequency subbands are used in this method as regression target values (Y_{mis}). Likewise, X_{obs} are used as features/predictors. Having set up this framework, we intended z to capture the interactions between the features of X_{mis} in order for them to generate a proper reconstruction of $Y_{missing}$ by maximising $P(Y_{Missing} \mid z; \theta_{dec})$ updating θ_{dec} (decoder's parameters). In other words, maximizing the likelihood of the training set under the generative model to being able to reproduce it. It is also crucial to update the encoder's parameters θ_{enc} in order to learn favourable latent space variables under $P(z \mid X_{obs})$.

4.4.2.1 Mathematical intuition of Variational Autoencoders (VAEs)

As we got inspired by VAEs, in this point we detail the mathematical notion of these generative models that enable the learning of our implemented DL architecture by using Variational Inference (VI) techniques. The assumption in VAE is that z can be drawn from $\mathcal{N}(0,I)$. This is argued by the possibility of reconstructing a dataset X departing from some latent independent normally distributed variables z undergoing a powerful enough function approximator $f(z;\theta)$. The following explanation is based in [37], [30], [38]. In order to maximise Equation 4.27 we need to account only for the probabilities P(X|z) that contribute significantly to estimate P(X). For that reason, a function P(z|X) is considered to take a value X and generate a distribution over z values that are more likely to generate a proper reconstructed X, and hopefully the space of z is smaller and easier to explore that the one under P(z). To approximate this posterior P(z|X) we use a VI framework with q(z):

$$q(z) = \arg\min_{q} KL(q(z) \mid P(z \mid x)). \tag{4.28}$$

KL is the Kullback-Leibler between q(z) and $P(z \mid x)$, let it be written as:

$$KL(q(z)||p(z|x)) = E_{q(z)}[\log q(z)] - E_{q(z)}[\log p(z,x)] + \log p(x)$$
(4.29)

Rearranging Equation 4.29, it can be noted that the KL element is by definition equal or larger than zero, therefore maximising log p(x) is to be attained by the Evidence Lower

Bound (ELBO). This is made explicit in Equation 4.30.

$$\begin{split} \log p(x) &= KL(q(z)||p(z|x)) - E_{q(z)}[\log q(z)] + E_{q(z)}[\log p(z,x)] \\ &\geqslant -E_{q(z)}[\log q(z)] + E_{q(z)}[\log p(z,x)] \\ &= -KL(q(z)||p(z)) + E_{q(z)}[\log p(x|z)] = ELBO \\ &= E_{q(z)} \left[log \frac{p(x,z)}{q(z)} \right] \end{split} \tag{4.30}$$

Note we introduced the distribution q(z|x) := p(z|x) such that it represents the encoder NN, that implicitly models the latent space conditioned on the data, for then reconstructing it by $p(x \mid z)$ as a decoder. Thus, when aiming to maximize the *ELBO* term, we allow for the VAE's parameter optimization as it implies increasing the marginal p(x) or decreasing the KL divergence.

The encoder distribution $p(z \mid x)$ and the prior for latent variables p(z) can be deemed to be Gaussian and approximated by the correspondent NNs. Note that in the following Equation 4.31 σ^2 and μ depend explicitly on result of the operations between the x fed to the encoder and encoder parameters θ , considered in a normal distribution that yield z by sampling.

$$q(z|x) = \mathcal{N}(z; \mu(x;\theta), I(\sigma^2(x;\theta))),$$

$$p(z) = N(z; 0, I).$$
(4.31)

4.4.2.2 Reparametrization trick

This setup allows us to perform feed-forward data passes trough the network as we can sample and average z from $q(z \mid x)$ as well as the gradient in ELBO loss. However, we still need to back-propagate the gradient for the weight update in training, which conflicts with $z \sim q(z \mid x)$ and $E_{q(z)}[\log q(z)]$, which is not continuous and lacking of gradient as they come from sampling layers. To work this around, what is performed is the reparametrization trick [30], that consists in embedding the sampling to an input layer, see Figure 4.3. The mean and covariance of $q(z \mid x)$, $\mu(x)$ and $\sigma^2(x)$, are used in combination with another sample $\varepsilon_i \sim \mathcal{N}(0,1)$ to draw our latent variable $z_i = g(x, \varepsilon_i) = \mu_i(x) + \sigma_i^2(x) * \varepsilon_i$. The yielded reparametrized loss expression that is a deterministic function and thus, across a sample of $x_m \in \mathbb{R}^R$ the loss function reads as follows:

$$\mathcal{L}_{VAE}(\theta) = KL(q(z|x_m)|p(z)) - \frac{1}{L} \sum_{l=1}^{L} \ln p(x_m|z_{l,m}),$$
(4.32)

where L is the number of Monte Carlo samples to average over the term $E_{q(z|x)}[\log p(x|z)]$.

4.4.2.3 Inside the Kullback-Leibler divergence

According to Kullback [39], the KL divergence between to Gaussians is analytically feasible. Consider we have under diagonal covariance Gaussians of dimension Q:

$$KL(\mathcal{N}(z_i; \mu_{1,i}, \sigma_{1,i}^2) || \mathcal{N}(z_i; \mu_{2,i}, \sigma_{2,i}^2)) = \frac{1}{2} \ln \frac{\sigma_{2,i}}{\sigma_{1,i}} + \frac{\sigma_{1,i}^2}{2\sigma_{2,i}^2} + \frac{(\mu_{1,i} - \mu_{2,i})^2}{2\sigma_{2,i}^2} - \frac{1}{2}.$$
(4.33)

Let the second Gaussian have $\mu_{2,i}=0, \sigma_{2,i}^2=1$ and to abbreviate $\sigma_i^2:=\sigma_{1,i}^2, \mu_i^2:=\mu_{1,i}^2$. Then we yield the loss term entailing the KL divergence:

$$KL(p(z_i|x)|p(z_i)) = -\frac{1}{2}\ln\sigma_i + \frac{1}{2}\sigma_i^2 + \frac{1}{2}\mu_i^2 - \frac{1}{2}.$$
 (4.34)

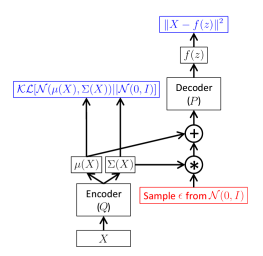


Figure 4.3: Variational autoencoder with parametrization trick (obtained from [37]).

After that, we need the negative log-likelihood, the so-called reconstruction loss $-\log p(x|z) = -\sum_{i=1}^R \log p(x_i|z)$. Let the mentioned R be the output dimension of the decoder, that we deem Gaussian as per assumption $p(x_i|z) = \mathcal{N}(x_i; \mu_i(z), \sigma^2(z))$. Mind that $\mu_i(z)$ is the output of the decoded NN. In practise, what it is forced to be outputted by the decoder instead of σ^2 is $l_i(x) := \ln \sigma^2(x)$ to force $\sigma^2(x) = \exp(l_i(x))$ to be positive. Following the logic of a continuous Gaussian output, the log-likelihood leads to the sum-of-squared error, leveraged as reconstruction loss, analogue to 4.26.



Experimental Set-up

This section is meant to be a thorough explanation on how the experiments with our already explained models were empirically developed (Sections 5.2 and 5.3). Prior to that, in Section 5.1 we describe the software used for each model employed in the thesis.

5.1 Software

5.1.1 Software - PPCA/BPCA

The programming language used to perform the PPCA's set of experiments is R [40], known for its wide-spread usage in statistical and probabilistic modeling. The main library used in these section is pcaMethods, leveraged to extract the PPCs and BPCs of our provided datasets. We also used the library MASS [41] for some statistical modeling. The fitting algorithms for both BPCA and PPCA implemented in pcaMethods are iterative and grow $O(n^3)$ due to the matrix inversions involved.

5.1.2 Software - VED

The programming language used to perform the VED's set of experiments is Python [42]. The main libraries used in these section are Keras [43] and TensorFlow [44], in order to build the VED architecture. We also used Scikit-Learn [45], Pandas [46] and Numpy [47] to transform and normalize the data. The plots in these experiments where provided by the package Matplotlib [48]. The Scipy.Stats module [49] was used in order to apply statistical modeling to the error of prediction (vector of $VEC(y_{predicted}) - VEC(y_{true})$, where VEC() stands for the function that flattens a matrix into a vector).

5.2 PPCA - Experiments

In this section we describe the steps carried out for the PPCA training and assessment.

5.2.1 Number of PPCs choice

The function ppca, was used in the training split of the different datasets in order to obtain a set of values: loadings (W, the probabilistic principal components), scores (Z), center $(\mu, \text{ the historic mean of every feature})$, R2 (% of explained variance), R2cum (% of cumulative explained variance).

Afterwards, a plot of the R2 was performed in order to decide the optimal number of PPCs (denoted by nPPCs). In this datasets, the maximum computation supported was 40 PPCs, which is deemed as enough because in all cases the % of accumulated variance stagnated from some prior point onwards.

Looking at the graphs in Figure 5.1, we used the heuristics of picking out the nPPCS that is prior to the stagnating. There, adding more complex to the model seems not explain more variance, thus leading to possible overfitting.

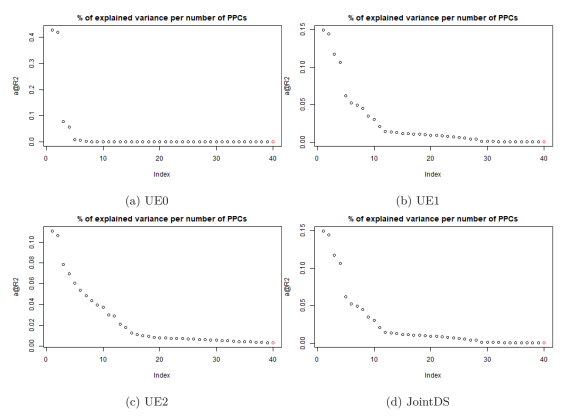


Figure 5.1: % of explained variance per number of PPCs (Own elaboration).

Therefore, the number of PPCs used in the model for each dataset are these: nPPC(UE0)=8, nPPC(UE1)=10, nPPC(UE2)=15, nPPC(JointDS)=12.

5.2.2 PPCA Extraction

Once the optimal nPPCs is chosen, we performed the same parameter extraction as in Section 5.2.1. If we want to apply the predictive function of Figure 4.11 we still miss the $I\sigma^2$ term. We estimated this term by performing $I\hat{\sigma}^2 = I \cdot VAR(\epsilon) = I \cdot VAR(VEC(WZ^T - X_{train}))$.

5.2.3 Prediction - PPCA

Leveraging the predictive function in Equation 4.11, we predicted inside each TTI of the validation set. We used the information available in X_{obs} to extrapolate on the $X_{missing}$

MPCs. The results are available in the PPCA Results Section 6.2. In order to take W_{mis} and W_{obs} , we extracted the rows in W that corresponded to the mis or obs indexes in each TTI. An analogue procedure was carried out for vectors μ_{mis} and μ_{obs} .

We also provided Confidence Intervals on this prediction matter by using the predictive function for the covariance in Equation 4.12 and applying the empirical rule $\hat{y}_i \pm 2\sqrt{\hat{\Sigma}_i}$ (where \hat{y}_i is the multivariate prediction for the TTI/observation i).

5.2.3.1 Cholesky decomposition

In the prediction step, the inversion matrix uses Cholesky decomposition [50]. This is done to deal with the huge dimensions of the covariance matrix to be inverted, which is also highly correlated. This is implemented for the expressions in Equations 4.11 and 4.12, in the form $\mathtt{chol2inv(chol(A))}$. This bit of code calculates the inverse of a symmetric, positive definite square matrix from its Cholesky decomposition. These technique follows the reasoning in Equations 5.1, 5.2, 5.3. Let A be the matrix to invert whereas $LL^T = A$ represents the Cholesky decomposition. Note that in Equation 5.1, L is a lower-triangular matrix, this implies that calculating its inverse is faster and more numerically/stable that using the inbuilt $\mathtt{solve()}$ function of R.

$$Chol(A) = L^{T} \Rightarrow A = LL^{T}$$

$$LL^{T}x = I$$
(5.1)

Let us introduce the intermediate variable $v := L^T x$. The main idea of inversion by decomposition is the performed in two steps presented in Equation 5.2 and 5.3:

$$Lv = I (5.2)$$

Let us first solve for v and then do:

$$Chol2inv(Chol(X)) \Rightarrow L^T x = v.$$
 (5.3)

Finally, solving for x in 5.3 we obtain $x = A^{-1}$.

5.2.4 BPCA - Experiments

The processes carried out in the BPCA experiments where analogue to the ones in the PPCA. We used the function bpca instead of ppca and the Bayesian Principal Components are extracted in an analogue way as described in Section 5.2. We used of the number of components chosen in Selection 5.2.

5.2.5 Assessment - PPCA

The metrics used to assess the regression performance in the validation set are the following:

$$MSE = \frac{1}{D} \sum_{i=1}^{D} (x_{mis} - \hat{x}_{mis})^2$$
 (5.4)

$$MAE = \frac{1}{D} \sum_{i=1}^{D} |x_{mis} - \hat{x}_{mis}|$$
 (5.5)

$$NRMSE = \sqrt{\frac{1}{D} \sum_{i=1}^{D} \frac{(x_{mis_i} - \hat{x}_{mis_i})^2}{mean(x_{mis}^2)}}.$$
 (5.6)

Qualitative plots of the fit of this model are also provided are also provided in Section 6.2.

5.3 VED - Experiments

5.3.1 Architecture Design

Our VED architectural design is based in the one described in the Keras documentation [51] and the already mentioned work of [28]. We opted for the concatenation of a encoder-decoder whose architecture varies for each dataset. We display the architecture choice for Dataset UEO, described in Tables 5.1 and 5.2. The rest architecture parameters for the other datasets are in the Section A of the Appendix. In this tables the "Output Shape" column means the number of hidden units for the layer, whereas "Param #" corresponds to the number of parameters to be trained in that layer (# Parameters = [# input weights of the layer + 1 bias weight] × # hidden units in the output).

The encoder and decoder were concatenated with a sampling layer in-between that has been coded as in Listing 5.1. This sampling layer is the one that provides the probabilistic/variational inference quality to the Encoder-Decoder. Note that in this bit of code we express the property explained in Section 4.4.2.3 of forcing $\log \sigma^2$. In general, this sampling layer yields a sample from a Normal distribution of [NxM]. N stands for latent dimensions and M for the dimensions provided by the batch size. This is possible by means of the parametrization trick. Please see Figure 5.2 where the diagram of our architecture is visually shown.

5.3.1.1 Heuristics in the Architectural Design

As said, we departed from the design in [28] to base our architecture in some optimal design decision that has been proved to work well in the literature. However, it was necessary to make changes on the architecture because the NN was not learning properly. The number of dense layers was reduced backing this decision up by the loss plots during the training phase (shown in Section 6.3), because the more complex the architecture was getting, the more unstable the learning became. The latent space dimension was also tweaked heuristically according to the same criteria, stable and convergent learning.

5.3.2 Training Customisation

For the total loss, we incorporated the KL-divergence in addition to the mean-squared error obtained by comparing X vs Y of every batch. We used implemented Early Stopping as a

Table 5.1: Model summary for the Encoder NN (UE0).

Model: Encoder

Layer (type)	Output Shape	Param	Connected to
input_1 (InputLayer)	(None, 618)	0	[]
dense_1 (Dense)	(None, 309)	191,271	['input_1[0][0]']
z_mean (Dense)	(None, 100)	31,000	['dense_1[0][0]']
z_log_var (Dense)	(None, 100)	31,000	['dense_1 [0][0]']
sampling_1 (Sampling)	(None, 100)	0	['z_mean[0][0]',
			$'z_log_var[0][0]']$

Total params: 253,271 Trainable params: 253,271 Non-trainable params: 0

Table 5.2: Model summary for Decoder NN (UE0).

Model: Decoder

Model. Decodel		
Layer (type)	Output Shape	Param #
input_1 (InputLayer)	(None, 100)	0
dense_2 (Dense)	(None, 50)	5,050
dense_3(Dense)	(None, 1,854)	94,554

Total params: 99,604 Trainable params: 99,604 Non-trainable params: 0

callback, although history plots of both the validation set loss and training set loss along the training for all epochs are provided. It would be useful for a testing subset assessment nevertheless.

The Batch Sizes for the VED vary accordingly to each dataset experiment heuristically. Again, the criteria was to ensure a smooth training that converged. We opted for Adam (explained in Section 4.4.1.3) as an adaptive stochastic gradient descent algorithm, and ReLU as an activation function (remember Equation 4.20). We implemented as well a hyperparameter to control the weight in the KL-divergence to the contribution in the total loss. We make it explicit it in Equation 5.7 as β but we finally set it to $\beta=1$ for all experiments as the training converged for both losses (MSE and KL) eventually.

$$\mathcal{L}_{\text{VAE}}(\theta) = \beta K L(q(z|x_m)|p(z)) - \frac{1}{L} \sum_{l=1}^{L} \ln p(x_m|z_{l,m})$$
(5.7)

5.3.3 Assessment - VED

For the assessment of the VED execution, we relied on the metrics presented in Section 5.2.5 so as we can fully compare model executions in the very same validation set. We also provide in Section 6.3 different plots akin to the ones related to the PPCA experiments.

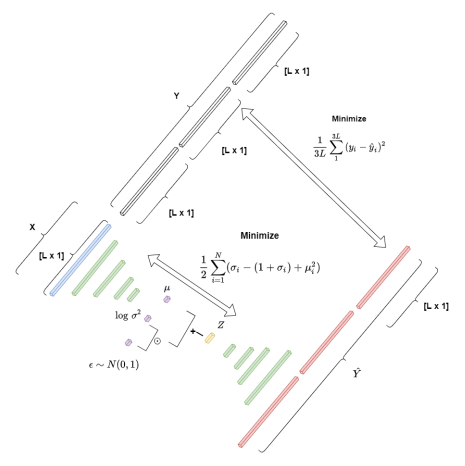


Figure 5.2: Architecture designed for the VED used in the thesis. The loss function to minimize corresponds to the KL-divergence and the one in the top-right corresponds to the MSE over the predictions against their background truth (Own elaboration).



Results and Discussion

This section presents the results of the PPCA, BPCA and VED models. It also contains further elaborations on these results. Herein we highlight the results that stand out further relating the outcome obtained with the underlying theory. We also discuss what is our intuition on what could happen if the results were undesirable. Moreover, in Section 6.1 we explain the assessment and exploration performed on the data that justifies some of the experimental/design decisions.

6.1 Preliminary Statistical Exploration of the Data

We inspected the data before carrying out the main experiments by using the MASS R package. We leveraged the functions **fitdistr** to tentatively fit Normal and Student-t distributions curves in order to know how the data could be roughly modeled. This function uses the Maximum Likelihood Estimation (MLE). The following subsections describe the result of this modeling to further obtain preliminary notions of how the data is distributed.

6.1.1 MLE Modeling on the Unmasked Training Set

We applied this MLE modeling to the unmasked training splits of the datasets. The curves of the Gaussian and the Student-t distributions are presented in Figures 6.1a and 6.1b for UE0 and UE1 datasets. The data in the histogram is clearly reflected not to be Gaussian (so was this reflection in the remaining datasets). As for this modeling, when the Gaussian curve is not able to capture all the data in the tails' support, it attempts to widen itself increasing the variance, hence trying to cover the most deviated data. In this way, it fails at enveloping the whole histogram (it does not envelope the central part where there is most of the density). In the case of the Student-t distribution, by tweaking the parameter of df (degrees of freedom that control the Kurtosis) the data gets modeled best. We assume that the rest of the data (the validation set) shares the same distribution as the training data.

6.1.2 Modeling the PPCA Reconstruction Error

As we have seen in Section 6.1.1, the data is not normally distributed. We extracted Probabilistic Principal Components (PPCs) and applied the reconstruction of the data through

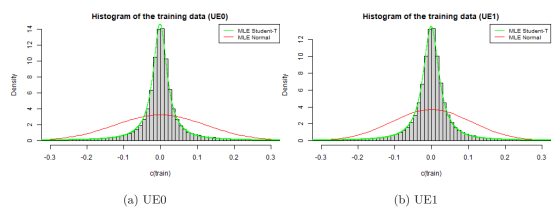


Figure 6.1: MLE modeling on the training subset (Own elaboration).

multiplying the obtained $W \times z^T$ (loadings times transposed scores). Then we subtracted the product from the training dataset to get the error: $\epsilon = X_{train} - W \times z^T$, as described in Section 4.1.

Hence, we overlaid the MLE curves on the histogram of this error ϵ . In the case of UE0 (Figure 6.2a), the error is considerably normal distributed, therefore we can assume that the 8 PPCs extracted capture the distribution as the reconstruction yields this $\epsilon \sim \mathcal{N}$. However, more density of probability is spread towards the tails in UE1 train dataset, Figure 6.2b. That reflects the inability of PPCA to preliminary capture the distribution of the data as the error cannot be approximated as Gaussian (the outcome in the remaining datasets resembles most to this UE1 result). We could tentatively address this problem to the fact that the CSI of some TTIs would not belong to the same coherence window anymore. It is not deterministically clear what the actual issue could be, yet accounting for the fact of having such thick tails we could consider the error falls more apart from $\mu=0$ because of the high variability in CSI over time.

As for UE1, we intuited that the PPCA predictive capability would not fare as well as in UE0 dataset due to the non-gaussianity of its reconstruction error ϵ and the intrinsic non-gaussianity of the data. We observed a similar behaviour in the modeling between UE1 and UE2.

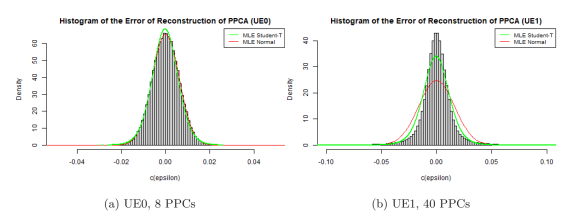


Figure 6.2: MLE modeling on the PPCA reconstruction error, training subset (Own elaboration).

6.1.3 Estimated Covariance Heatmap

Let us consider WW^T as an approximation of the covariance Σ taken out of UE0 dataset, obviating the term σ^2I . Note that W is the factor loadings extracted from the PPCA fitting process, remind Equation 4.2. We made use of the library Seriation [52] to obtain a heatmap of this covariance matrix. This library was used just for plotting purposes, no internal algorithm from Seriation was used, we just plotted WW^T with it. Mind the truncation trick used in Listing 6.1 to augment the resolution of the heatmap of Figure 6.3.

In this Figure, every pixel represents the correlation between each of the 2,472 of the effective unfolded UE0 dataset. We can observe there exists a pattern of squares. Each square in the diagonal represents the correlations within either the real or imaginary features belonging to the same Frequency SB. Mind that, for instance, the first column of pixels of the matrix correspond to the correlation of real part of the first MPC of the 1st SB $(\mathfrak{Re}(MPC_1^{SB_1}))$ with the rest of other features. The imaginary part of this 1st MPC $(\mathfrak{Im}(MPC_1^{SB_1}))$ shows its correlations versus all the other features in the 1st column of the second square. The off-diagonal squares represents correlations of either real or imaginary features against their conjugate counterpart in the same SB or against the real or imaginary features of other subbands. In Figure 6.4, two closeups of the Figure 6.3's heatmap are shown. The right heatmap is a slice of the correlations of two Frequency SBs, whereas the left image corresponds to the real features of the first SB. At square level, the higher correlations are in the diagonal (auto-correlation) and in the top-left section, meaning that there is a stronger correlation between the first features (MPCs) of each Frequency SB (on its own) and between the first MPCs belonging to different Frequency SBs.

When looking at the whole plot, there is a hardly visible unitary diagonal representing the auto-correlation of the features. There are not observed high correlations in the elements close to the diagonal that make it thicker. This could be originated because the most relevant correlations are between the first 25% (approximate) MPC of each SB (top-left corner of each square).

```
S <- pc_loadings %*% t(pc_loadings)

max = 2 * sd(S)

S[which(S>=max)] = max
S[which(S<=-max)] = -max

Listing 6.1: Truncation of WW<sup>T</sup> values, (R Code)
```

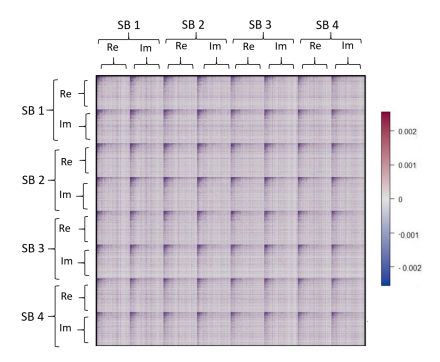


Figure 6.3: Heatmap of the WW^T , [2,472 x 2,472] (Own elaboration).

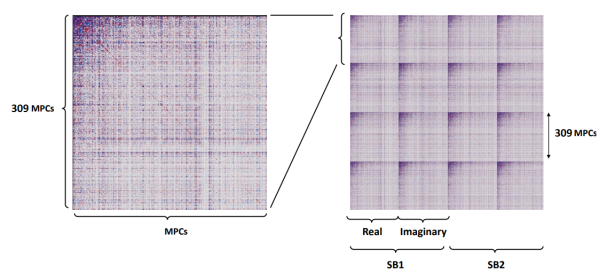


Figure 6.4: Close-up of Approximated Covariance Heatmap (Own elaboration)
Left, [309 x309] heatmap of the Real part of the MPCs belonging to the 1st Frequency SB.
Right, correlations of the features belonging to the 2 first SBs, Real and Imaginary parts highlighted.

6.2 Results - PPCA & BPCA

By following the described prediction algorithm in Section 5.2, we show in Table 6.1 the results of the prediction in the validation set of each dataset employed. As a visual support of these prediction outcomes, we attach several plots that reflect how the model executions developed.

		Dataset			
Experiment	Error Metric	UE0	UE1	UE2	JointDS
	MAE	0.0276	0.0513	0.0720	0.0664
PPCA	RMSE	0.0545	0.1062	0.2417	0.1573
	MSE	0.0029	0.0112	0.0584	0.0247
	MAE	0.0276	0.0514	0.0996	0.0665
BPCA	RMSE	0.0545	0.1064	0.5892	0.1568
	MSE	0.0029	0.0113	0.3471	0.0245
nP	PCs	8	10	15	12

Table 6.1: Error Metrics of PPCA/BPCA Experiments.

In Figure 6.5, it is shown how the predictions are scattered versus the corresponding background truth values for the PPCA and BPCA learnt models in all of the four datasets. In this plot, every (x_i, y_i) point correspond to $(y_{true_i}, y_{predicted_i})$. Thus, the thinner the cloud of points, the better the prediction. Likewise, a slope close to 45° with respect to the horizontal implies a better overall prediction.

Moreover, in Figure 6.6 we show how the predictions are visually presented over all the features across one time sample (one TTI) for the PPCA model in datasets UE0 and UE2 (remember Figure 3.4 in Section 3). In Subfigure 6.6a we can see that the image presents considerably less distant predicted values (red) to the background truth than in Subfigure 6.6b. This is reflected in the results of Table 6.1 if extrapolated for all the dataset. Note also that there are black dots in Subfigure 6.6b quite deviated from the main cloud of points.

To visually asses how the predictions behave over the intended support, we sorted all the background truth values of the validation subset by amplitude and overlaid the corresponding predictions (see Figure 6.7). In this figure we show the sorted predictions of datasets UE0 and UE1. Note that the predictions of UE0's validation split seem to capture more properly the range of values. UE2 and JointDS datasets incur the same behaviour as the shown UE1 in Figure 6.7b, see Appendix B for related figures.

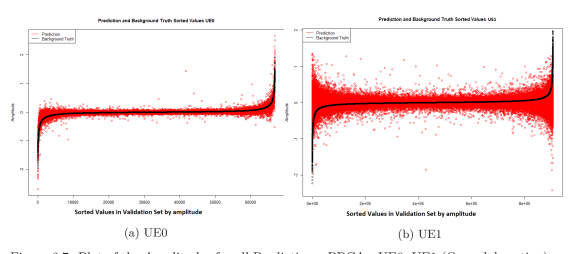


Figure 6.7: Plot of the Amplitudes for all Predictions. PPCA - UE0, UE1 (Own elaboration). Note the black dots represent the actual observed Multipath Components (Re o Im part), whereas the red correspond to the predictions.

We also wanted to assess some measures of uncertainty in the predictions. Therefore, we display in the Figure 6.8 how the confidence intervals look in the predicted, sorted values of one single TTI. There, it is obvious that in the case of the UE0 dataset, the predictions are more certain than in UE1. Taking a closer look to Subfigure 6.8a, we observe wider CI bands

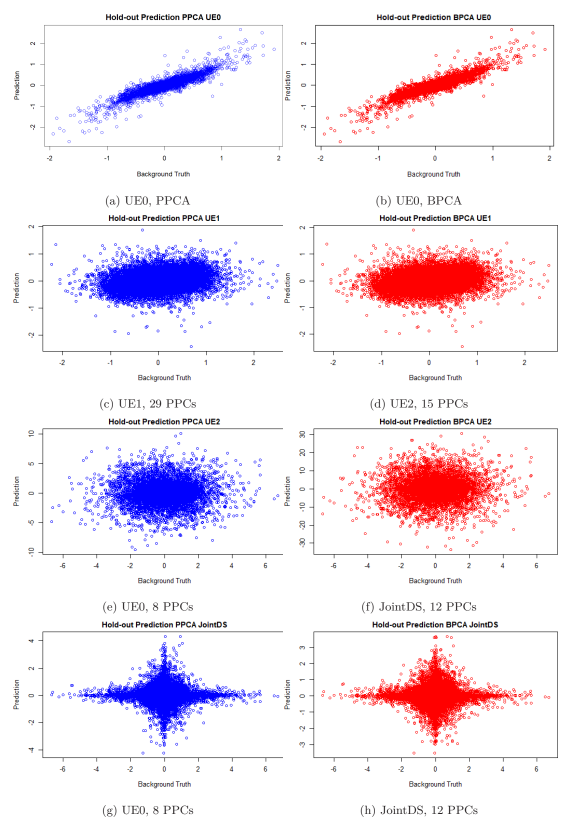


Figure 6.5: Validation subset prediction plot for the missing values (PPCA) (Own elaboration). \hat{y} On the vertical axis versus y_{true} in the horizontal.

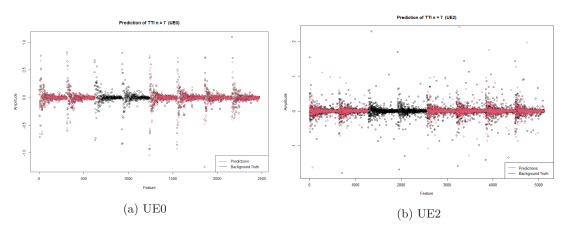


Figure 6.6: Plot of the Amplitudes for one TTI. PPCA - UE0, UE2 (Own elaboration). Note the black dots represent the actual observed Multipath Components, whereas the red correspond to the predictions. The segment of black values with no overlaid red is the observed Frequency Subband.

than in Subfigure 6.8a. Both Subfigures, though, present more uncertainty the closer they are to their mean, which is near to the 0 amplitude value. The fact that the prediction of the model is more certain in UE0 matches what is already explained about Figure 6.6, less distant values compared to the background truth. See more related Figures for the remaining datasets in Appendix B.

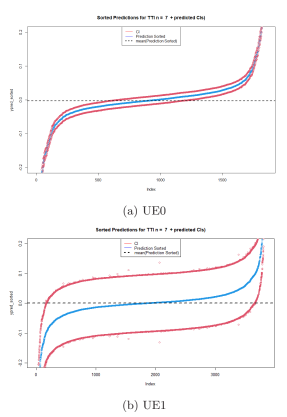


Figure 6.8: Confidence Intervals plot in the prediction of a $TTI.\ PPCA$ - $UE0,\ UE1.\ (Own\ elaboration).$

These plot shows the sorted redicted values for 1 TTI along with the Confidence Intervals overlaid.

Finally, we took the predicted errors $(VEC(y_{true}) - VEC(y_{prediction}))$ for each dataset's validation subset in a 1:1 fashion and flattened them into their respective 1-dimensional error vectors. Later on, we applied the same MLE modeling we performed in Sections 6.1.1 and 6.1.2 to these flattened vectors. We show these modeling outputs in Figure 6.9. Note the modeled red MLE-fitted Gaussian curve in Subfigure 6.9a is the one that seems to be capturing the data in the histogram the most. In overall, all the plots in all these Subfigures look like Student-T-distributed rather than Gaussian distributed. This could be alleged to the inability of PPCA to capture the dependencies between the features by linear-gaussian modeling.

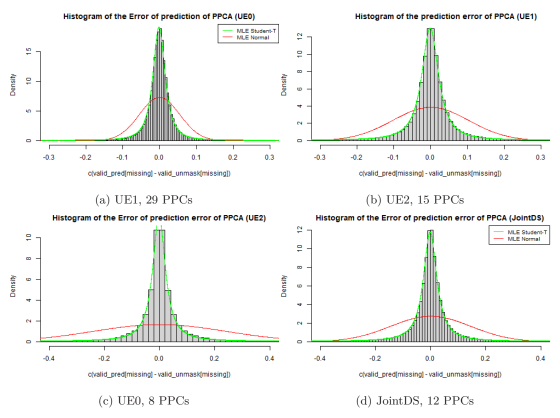


Figure 6.9: MLE modeling on the PPCA prediction error, validation subset (Own source).

6.2.1 Discussion on the PPCA and BPCA Results

Once the model was fitted and we further carried out the predictions and the statistical modelings, we could observe that in overall the PPCA model attained better performance than BPCA. The fact that the BPCA slightly underperformed the PPCA could be because the library implementation designed by Oba [24] is intended for missing value estimation rather than a hold-out training/prediction approach. As orthogonality is not forced in the fitting of the factor loadings, that may suppose that the new coordinate system defined for them does not capture the greatest variance for the scalar projections of the data as optimally as PPCA does. This method is alleged to perform better when the missing values are embedded in the datapoints that make up the training set where the EM fitting is performed. That is to say, in a value imputation approach. However, as seen in Figure 6.5, both methods present very similar predictive behaviours.

According to Figure 6.5, the best prediction is undoubtedly the one in UE0 dataset, see Subfigures 6.5a and 6.5b. It can be argued that in that dataset the predictions are more accurate (mind also that in Table 6.1 it presents the best results) due to the scarcity of TTIs

in comparison to the other datasets. Remember that the variation in the Channel becomes less linear and smooth the more we get far in time from the Coherence Window. As this interval is suppose to be around 80 TTI in the simulated datasets, we expect an overall decent performance in this dataset.

The distributions of the predictions of UE1 and UE2 (Subfigures 6.5c to 6.5f) do not show a cloud of points with such a good inclination. They also reflect thicker clouds of points. We could map the thickness of the clouds to the variance/uncertainty or instability in the predictions, since for similar background truth values the model is predicting a remarkably different range of values. The inclination is more intrinsically related to how good in average the model is able to predict properly for the whole support of values. Then, having a horizontal cloud of points means a poor fit of the model, as it is in average predicting the same for all the support of background truth values. We could blame this drastic deterioration on the prediction to more pronounced changes in the Channel over time, due to the large quantity of TTIs in comparison to the Coherence Window.

In the case of JointDS, Subfigures 6.5g and 6.5h, the prediction distribution seems to reflect a bivariate Student-T distribution. The fact this dataset is a merge out of UE1 and UE2 could have yielded some counterproductive dependencies in the process of covariance approximation of PPCA. In this Subfigures it can be clearly observed that the prediction is completely random for the amplitude mean of the background truth data (close to 0). Conversely, it predicts 0 for the rest of the support of background truth values.

In general, the less Gaussian-distributed was the reconstruction error of the datasets (see plots in Figure 6.2) the better the prediction was. We could relate this property to the Gaussian assumption of the PPCA/BPCA models in the data and the error. See how in Figure 6.9 the error is mostly Student-T-distributed instead of Gaussian. In the case of UEO (Subfigure 6.9a) the Gaussian curve attains to fit the data better than in the rest of the cases. This phenomenon reflects how the Gaussian assumption prevents the model to capture the data under the density of probability in the tails, which is large.

This uncertainty in the prediction in UE1, UE2 and JointDS datasets is also reflected in SubFigure 6.8b, which in comparison to the uncertainty reflected in UE0 (Subfigure 6.8a) is much larger for most of the values, visible at the width of the Confidence Intervals. However, the plots in Figure 6.7 reflect how for UE1 (b), the values that go away from the mean are not captured (it happens the same for UE2 and JointDS, see Appendix B). The model just increases the variance for those amplitudes in that range, failing to output predictions with the correct sign. It is not the case for UE0 (a), where these outliers in the dataset are better captured by the PPCA model. This is also reflected in Figure 6.6, where for UE2, it is clearly visible that beyond some threshold, the predictions fail at capturing the datapoints. In UE0 (b), conversely, we see that PPCA is closer to capture these points. The property of PPCA of departing from the historical mean of the target feature and then leveraging the predictive conditioned covariance could be a explanation for the goodness of the predictions in UE0.

We should also point out that, besides the increase in regards to TTIs of UE1, UE2 and JointDS with respect to UE0, there also exist an increase in the feature space, from 2472 effective features to 5120. Our practical intuition leads us to think that having ten times more datapoints than features might have also helped the model to perform a better learning of the underlying dependencies. It is also remarkable that having to predict for 75% of the values at each datapoint has been such a determinant limiting factor for the aim of learning a reliable regression model.

6.3 Results - VED

After having trained the models and further predicted for the validation splits' values, we show in Table 6.2 the results of such experiments for each dataset employed. As a visual support of these prediction outputs, we attach several plots that reflect how the model executions developed.

	Dataset			
Error Metric	UE0	UE1	UE2	JointDS
MAE	2.1730	11.801	16.7114	28.525
RMSE	0.7671	1.6105	2.6820	3.1247
MSE	0.5864	2.5927	7.2108	9.7785

Table 6.2: Error Metrics of VED Experiments.

The tracked losses in the training stages of the four architectures are arranged in Figure 6.10. There, we can observe that both training and validation losses converge for all the datasets. Similar to what we did in the PPCA/BPCA experiments, we visually assessed how the predictions are distributed against their respective background truth values (Figure 6.11). Again, analogue to the PPCA experiments, in Figure 6.12 we present how, for a single TTI, the predictions are arranged. It can be noted that there is an inability of the model when predicting for values over a threshold. Although, these bigger values are better captured in the UEO experiment (6.12a).

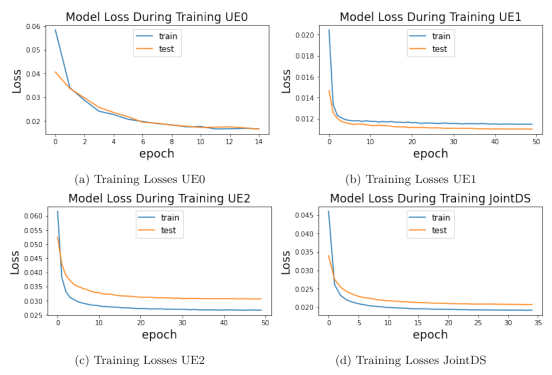


Figure 6.10: Training Losses for the NNs in all datasets (Training and Validation Splits) (Own elaboration).

We also performed the plots of all the sorted validation set's predicted values for UE0 and UE1 (Figure 6.13). Similar outcome was produced for the remaining datasets, see Appendix C. It is visible that the models have troubles to capture values larger than some range in positive and negative amplitudes. There also exists a great variance on the values predicted.

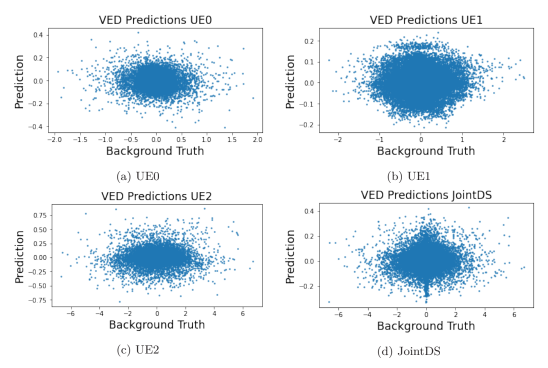


Figure 6.11: Validation subset prediction plot for the missing values (VED) (Own elaboration). \hat{y} on the vertical axis versus the \hat{y}_{true} in the horizontal.

The amplitudes unable to be reached are notably less present among the validation set, we could consider them to be outliers, even though the model fails at reaching them.

Finally, same modeling of the prediction error ($VEC(y_{predicted}) - VEC(y_{true})$) as in the PPCA/BPCA experiments is performed (this time with Scipy.Stats Python library), see Figure 6.14. Again the more similar to a Gaussian distribution looks to be UE0, but all of them are best fitted by a Student-t distribution.

6.3.1 Discussion on the VED Results

In this Section we proceed to carry out a general commentary on the results as we did in Section 6.2.1 for the baseline. Generally, the VED models obtained worse performances than the baseline, (around an order of magnitude of $\times 100$ in the case of the MSE). Ensuring a convergent and stable training was something we first struggled with when choosing the architecture hyperparameters (number of layers, type of activation functions, number of hidden units, dimension of latent space, loss function, etc.). Then, we tried to vary heuristically these hyperparameters in order for the output to try to capture the values of larger magnitude. In Figure 6.12 we see that the values beyond some threshold are not predicted properly. This is more accentuated in UE1 6.12b, where the span of prediction is narrower, even (same output for UE2 and JointDS, available in Appendix C).

The mentioned phenomenon is also pronounced in Figure 6.7, where we plotted for UE0 and UE1 the predictions of the validation set sorted by amplitude. As in the PPCA case, the model fails to capture these outliers. Note that we can see this fact reflected in the statistical modeling of the prediction error in Figure 6.14, where it is shown the incapability of capturing these extreme values in the thick tails for all datasets. Again, the Student-T distribution would be a more suitable distribution of this error.

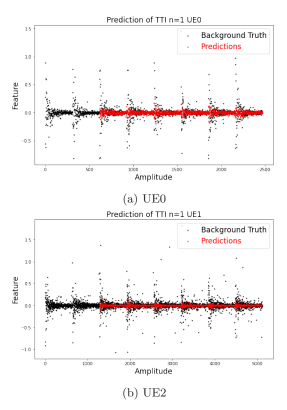


Figure 6.12: Plot of the Amplitudes for one TTI. VED - UE0, UE1 (Own elaboration). Note the black dots represent the actual observed Multipath Components, whereas the red correspond to the predictions. The segment of black values with no overlaid red is the observed Frequency Subband.

As a general note on the prediction quality, we presented scatter plots of the predictions versus the background truth values in Figure 6.11. All the clouds of points seem Gaussian-distributed, although the lack inclination and thickness reflect inaccuracy of the predictions.

In overall, it looks that the model has failed in capturing the data distribution, as well as in its main purpose, which was learning the non-linear dependencies of the features to increase the quality of the predictions in relation to the baseline. The squared pattern in the covariance heatmap of Figure 6.3 describes a strong dependencies of the first MPCs intra/inter-frequency subband. We wanted to exploit this scenario by performing the dimensionality reduction, thinking that this important dependencies would be kept and outweighed the irrelevant ones. We addressed the cross-band extrapolation problem by using a variational inference approach and relying in its generative distribution approximation power for a proper feature reconstruction. Generally, the model presents clear signs of not performing optimally since the output relies too much in predictions close to the training data mean, whose presence in the dataset is overly abundant (high kurtosis, due Student-T distributed modeled noise).

A possible downside of the model chosen could have been the Gaussian assumption of the loss function (MSE), that considers the output to be normally-distributed, despite the Statistical modeling performed reflects not to be the case. A likelihood function resulting from the Student-T distribution could be tried as an alternative loss function instead of the mean squared error, aiming to capture extreme values that fall more distant to the mean in the tails of the data distributions for all the datasets. In order to explain this density of values that are under the tails of the Student-t distributed data and prediction noise (error), we could look for potential covariates that could explain this behaviour. As the outliers of the dataset and the predicted values tend to be located at both tails of the distribution (i.e., extreme

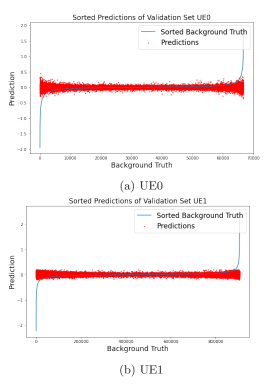


Figure 6.13: Plot of the Amplitudes for all Predictions. VED - UE0, UE1 (Own elaboration). Note the black dots represent the actual observed Multipath Components (Re o Im part), whereas the red correspond to the predictions.

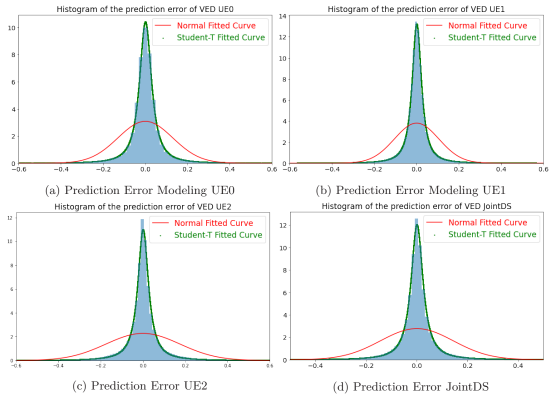


Figure 6.14: MLE modeling on the VED prediction error, validation subset (Own source).

negative and positive values), we intuit there exist certain pattern relative to the subband MPCs that could be exploited. The variation in the hyperparameter that regulates the weigh of the KL loss could also be explored in order to ensure that its contribution does not make the model get a better fit.

From a variance-bias trade-off perpective, it would be difficult to assess the performance based on the curves in Figure 6.10. Even though all training stages seem to converge, in 2 cases (UE2 and Joint DS datasets) the train loss ends under the test loss. In the case of UE0 the losses look like they converge to the same point whereas in UE1 the train loss is over the test loss. We will consider that generally the models underfit as the datapoints (TTI) are quite scarce in comparison to the features and there might be non-linear dependencies in the spatial domain that could have been not learned.

6.4 Discussion on an alternative data pre-processing configuration

Our intuition leads us to think that another pre-processing configuration on the data could be tried in future experiments. A smaller experiment was also run, where we used the polar plane coordinates for the MPCs instead of the Cartesian plane coordinates (as they are complex values). Remember that for the main work of the thesis we used the form z=a+bi (where we separated in columns the $a=\Re\mathfrak{e}(z)$ and the $b=\Im\mathfrak{m}(z)$). For the alternative configuration we used the form $z=\alpha/(\phi)$, where α is the modulus (amplitude) of one certain MPC in one certain TTI and ϕ is its phase. We separated angles and phases in columns, analogue to our pre-processing shown in Section 3.2. In the Figures 6.15 and 6.16, we show how the phase and the module of the first MPC for the four SBs evolves over time. Notes that the phase evolves in this discontinuous pattern (but step-wise linear) and the module in a sinusoidal way.

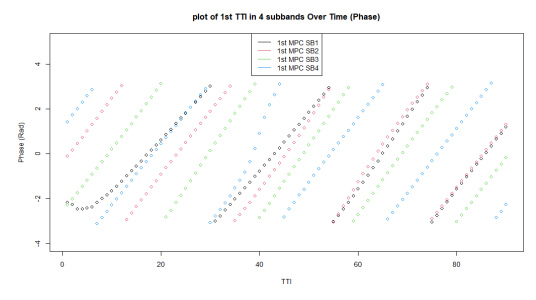


Figure 6.15: Plot of the evolution over time of the 1st MPC component in the 4 SBs, phase of the 1st MPC in the 4 SBs (UE0) (Own elaboration).

In Figure 6.17 we show the magnitude of the MPCs (their modules and phases) for the 4 SBs. Notice here the different magnitudes and ranges of the phases versus the modules. In Figure 6.18 we show a preliminary experiment of prediction over a TTI as we did in Section 6.2, using PPCA (analogue to Figure 6.12). Notice how the prediction over the modules is much better than in the phase. This could be due to an non-optimal normalization of the

Figure 6.16: Plot of the evolution over time of the 1st MPC component in the 4 SBs, module of the 1st MPC in the 4 SBs (UE0) (Own elaboration).

data. That is to say, as the phases and modules come from different distributions and have different ranges, the model had troubles in using a good predicting variance for the phase.

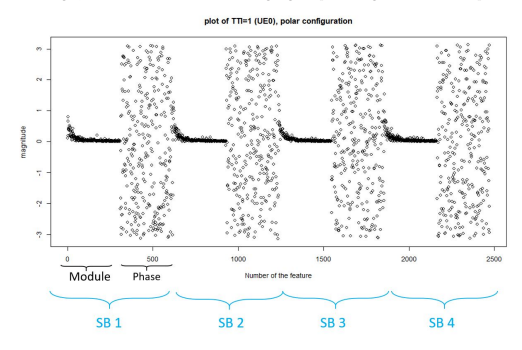


Figure 6.17: Plot of the MPCs in 1 TTI, modules and phases (UE0) (Own elaboration).

In spite of this experiment failing at predicting MPCs' phases, we intuit a promising potential in the module prediction that could be studied in future works. As for the phase prediction, it could be modeled per separate just accounting for the phase values across subbands and in the same subband across time.

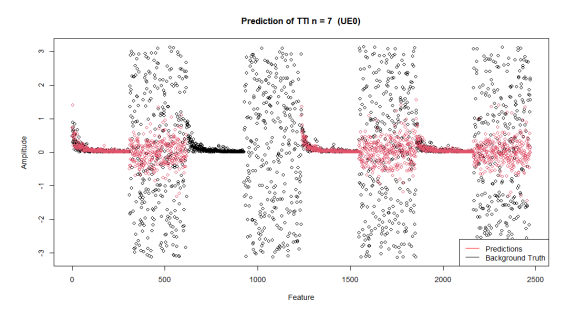


Figure 6.18: Plot of the PPCA prediction over the MPCs in 1 TTI, modules and phases (UE0). (Own elaboration).



Ethical Considerations

This thesis work is commissioned by The RD Center of Huawei Technologies Sweden AB in order to explore the Channel Prediction problem which is encompassed in the wireless communications field. There are several related considerations that could have an impact to society and the environment to some extent. The ultimate aim of this work is to increase the throughput capacity, data-rate, spectral efficiency and network capacity of wireless transmissions in MIMO scenario between users and base stations. To tackle this problem, we isolated the scenario into a SIMO approach, thus focusing in one UE and multiple antennas to try to revamp the communication transmission's quality.

For the safety of the environment and individuals, security policy standards that rule a fair use of the frequency bandwidth have to be followed depending on the technology used, such as 5G. Overlapping an unintended frequency bandwidth could occasion harmful troubles for the safety of other individuals and organizations. That is because this fact could compromise somenone else's communication due to interference. Another issue to comply with is the quality standards for a reliable communication that one has to aim for, such as the Regulation (EU) 2015/2120 of the European Parliament [...] towards electronic communications networks and services and Regulation (EU) [53].

8 Conclusion

It has been possible to create a method of cross-band channel prediction and further evaluate it with statistical metrics as well as with visual assessment. It turned out that the experiments conducted in UE0 were the most satisfactory ones. The most revealing assessment was the visual one we carried out by plotting the predictions along individual TTIs and over all the validation dataset. Through this visual, qualitative evaluation, we could address the good quality of this results to a successful covariance matrix approximation through dimensionality reduction by means of PPCA. This could be explained because all the TTIs might be forming part (approximately) of the same Coherence Window. Thus, linear dependencies between MPCs could be leveraged for the prediction of missing ones by using the observed information for each TTI.

In the case of the VED, the results were not so satisfactory, showing worse prediction results. We could address this problem to the underfitting occurred during the training, where the VED was reflected not to have captured the non-linear relations between the features for a proper extrapolation. Therefore, we propose some further work that could circumvent this event. A tweak that could be performed to the VED architecture would be to try a loss function of another nature, like the likelihood function of a Student-t distribution. Also, we could look for covariates that could explain the behaviour of the noise not captured (present in the tails of the Student-t modeled distribution). This would assume the data distribution not to be Gaussian, aiming to capture the outliers more effectively. Likewise, this shift of assumption might be worth to be assessed in the sampling layer, switching to a Student-t distribution in the bottleneck of the VED where the variational inference takes place.

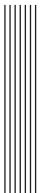
On another note, further regularization strategies should be tried, e.g., dropout. In this line, further hyperparameter searching could be implemented at the level of VED architecture: hidden units and layers, dimension of the latent space, activation function, etc.

On the evaluation matter, the assessment of the prediction's quality could be revamped by adding posterior stages of signal reconstruction. This would allow to use intrinsic wireless communications performance metrics such as beamforming gain, spectral efficiency and Signal-to-Noise ratio, that were not use in the thesis due to time and background constraints. These metrics would use the reconstructed H matrices instead of the MPC, allowing to compare the results directly to literature benchmarking and state-of-the-art.

Concerning the inter-TTI prediction purpose, due to the complexity of implementing a framework that combine feature-wise dependencies with time-domain ones, it has been an unfeasible goal that we must leave for a future research track. These approach could be worked out by the use of LSTMs as hidden layers in the VED, using different order p of previous TTIs involved in the time dependencies in the learning.

In matters of the data pre-processing, we would want to highlight that further experiments could be performed by using the Multipath Components data in the polar plane instead of in the Cartesian one. As we observed, there are strong patterns when looking at all the features in one time sample. This is more favourable when accounting for the module rather than in the phase. As for the phase, a more profitable pattern is observed over time for the MPCs independently (columns over time).

Finally, reproducing the experiments with a larger database in terms of TTIs (in the order of tens of thousands) could fare better with the currently implemented models, as it is usually a desirable method to prevent underfitting.



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Appendix - Architecture Parameter Design Tables

Table A.1: Model summary for the Encoder NN (UE1).

Model:	Encoder

Layer (type)	Output Shape	Param	Connected to
input_1 (InputLayer)	(None, 1280)	0	[]
dense_1	(None, 309)	395,829	['input_1[0][0]']
z_mean (Dense)	(None, 50)	15,500	['dense_1 [0][0]']
z_log_var (Dense)	(None, 50)	15,500	['dense_1 [0][0]']
sampling_1 (Sampling)	(None, 50)	0	['z_mean[0][0]',
			$'z_log_var[0][0]']$

Total params: 426,829 Trainable params: 426,829 Non-trainable params: 0

Table A.2: Model summary for Decoder NN (UE1).

Model: Decoder

Layer (type)	Output Shape	Param #
input_1 (InputLayer)	(None, 50)	0
dense_2 (Dense)	(None, 50)	2,550
dense_3 (Dense)	(None, $3,840$)	195,840

Total params: 198,390 Trainable params: 198,390 Non-trainable params: 0

Table A.3: Model summary for the Encoder NN (UE2).

Model: Encoder

Layer (type)	Output Shape	Param	Connected to
input_1 (InputLayer)	(None, 1,280)	0	[]
dense_1	(None, 350)	448,350	['input_1[0][0]']
z_mean (Dense)	(None, 100)	35,100	['dense_1[0][0]']
z_log_var (Dense)	(None, 100)	35,100	['dense_1[0][0]']
sampling_1 (Sampling)	(None, 100)	0	['z_mean[0][0]',
			$'z_log_var[0][0]']$

Total params: 518,550 Trainable params: 518,550 Non-trainable params: 0

Table A.4: Model summary for Decoder NN (UE2).

Model: Decoder

Layer (type)	Output Shape	Param #
input (InputLayer)	(None, 100)	0
dense_2 (Dense)	(None, 300)	30,300
dense_3 (Dense)	(None, 3,840)	1,155,840

Total params: 1,186,140 Trainable params: 1,186,140 Non-trainable params: 0

Table A.5: Model summary for the Encoder NN (JointDS).

Model: Encoder

Layer (type)	Output Shape	Param	Connected to
input_1 (InputLayer)	(None, 1,280)	0	[]
dense_1	(None, 309)	395,829	['input_1[0][0]']
z_mean (Dense)	(None, 100)	31,000	['dense_1[0][0]']
z_log_var (Dense)	(None, 100)	31,000	$['dense_1[0][0]']$
sampling_1 (Sampling)	(None, 100)	0	['z_mean[0][0]',
			$'z_log_var[0][0]']$

Total params: 457,829 Trainable params: 457,829 Non-trainable params: 0

Table A.6: Model summary for Decoder NN (JointDS).

Model: Decoder

Layer (type)	Output Shape	Param #
input (InputLayer)	(None, 100)	0
dense (Dense)	(None, 300)	30,300
dense (Dense)	(None, $3,840$)	1,155,840

Total params: 1,186,140 Trainable params: 1,186,140 Non-trainable params: 0



Appendix - PPCA Results Plots

Prediction and Background Truth Sorted Values (UE2 PPCA)

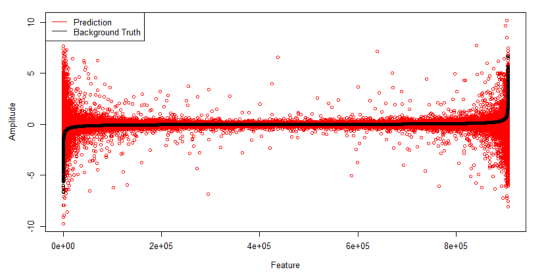


Figure B.1: Plot of the amplitudes for all predictions, PPCA - UE2 (Own elaboration) Plot of the amplitudes for all predictions. Note the black dots represent the actual observed Multipath Components (Re o Im part), whereas the red correspond to the predictions.

Prediction and Background Truth Sorted Values (JointDS PPCA)

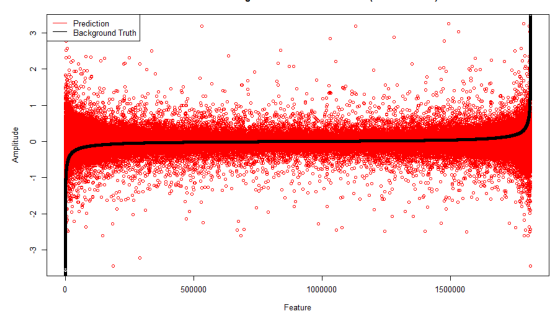


Figure B.2: Plot of the amplitudes for all predictions, PPCA - JointDS (Own elaboration) Plot of the amplitudes for all predictions. Note the black dots represent the actual observed Multipath Components (Re o Im part), whereas the red correspond to the predictions.

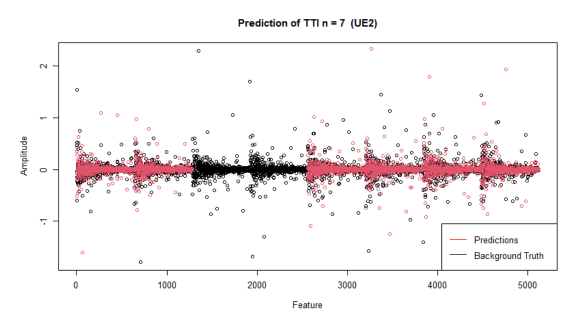


Figure B.3: Plot of the amplitudes for one TTI, PPCA - UE2 (Own elaboration)

Note the black dots represent the actual observed Multipath Components (Re or Im part),
whereas the red correspond to the predictions. The segment of black values with no overlaid
red is the observed Frequency Subband.

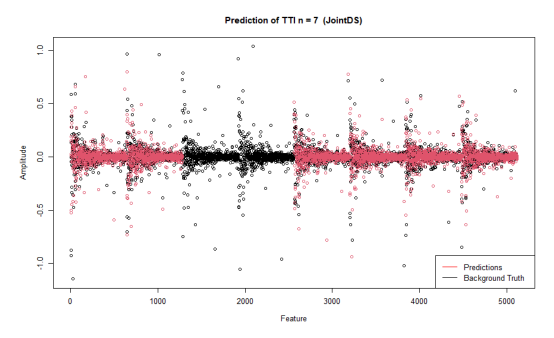


Figure B.4: Plot of the amplitudes for one TTI, PPCA - JointDS (Own elaboration) Note the black dots represent the actual observed Multipath Components (Re or Im part), whereas the red correspond to the predictions. The segment of black values with no overlaid red is the observed Frequency Subband.

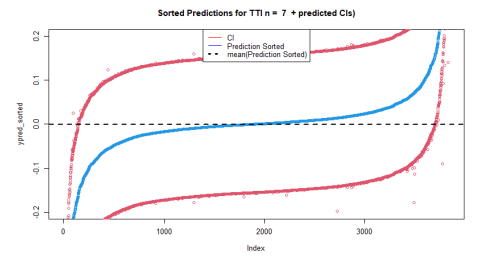


Figure B.5: This plot shows the sorted predicted values for 1 TTI along with the Confidence Intervals overlaid, UE2 (PPCA) (Own elaboration).



Appendix - VED Results Plots

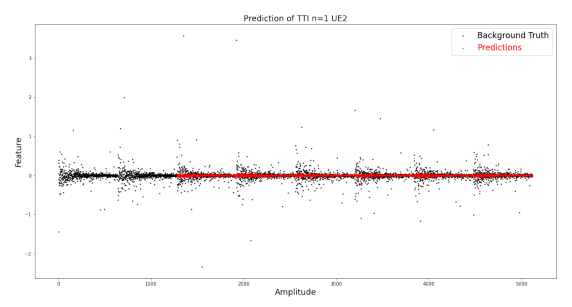


Figure C.1: Plot of the amplitudes for one TTI, VED - UE2 (Own elaboration)

Note the black dots represent the actual observed Multipath Components (Re or Im part),
whereas the red correspond to the predictions. The segment of black values with no overlaid
red is the observed Frequency Subband.

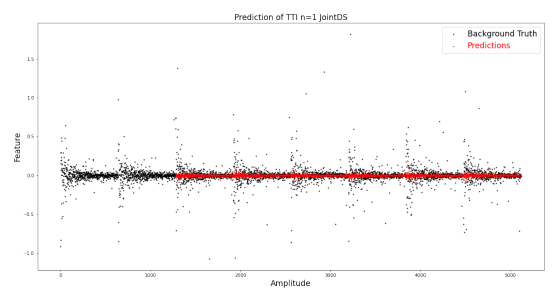


Figure C.2: Plot of the amplitudes for one TTI, VED - JointDS (Own elaboration)

Note the black dots represent the actual observed Multipath Components (Re or Im part),
whereas the red correspond to the predictions. The segment of black values with no overlaid
red is the observed Frequency Subband.

Prediction and Background Truth Sorted Values (UE2 PPCA)

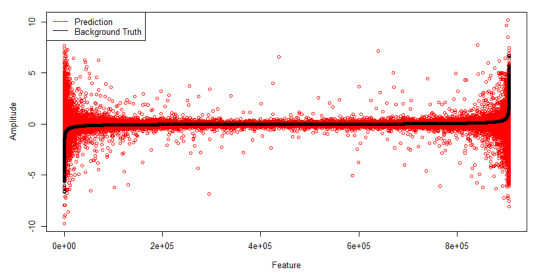


Figure C.3: Plot of the amplitudes for all predictions, VED - UE2 (Own elaboration)
Plot of the amplitudes for all predictions. Note the black dots represent the actual observed Multipath Components (Re o Im part), whereas the red correspond to the predictions.

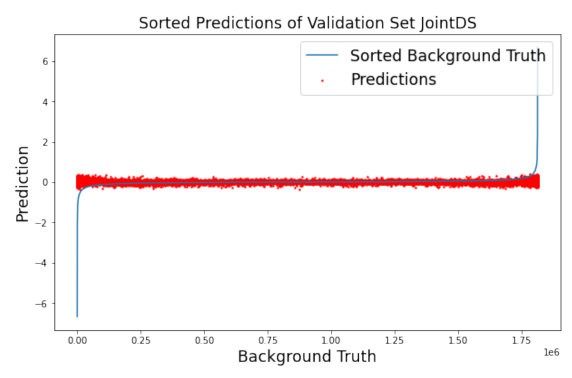


Figure C.4: Plot of the amplitudes for all predictions, VED - JointDS (Own elaboration) Plot of the amplitudes for all predictions. Note the black dots represent the actual observed Multipath Components (Re o Im part), whereas the red correspond to the predictions.