Complex-Valued Neural Networks With Nonparametric Activation Functions

Simone Scardapane ©, Steven Van Vaerenbergh, Senior Member, IEEE, Amir Hussain ©, Senior Member, IEEE, and Aurelio Uncini, Member, IEEE

Abstract—Complex-valued neural networks (CVNNs) are a powerful modeling tool for domains where data can be naturally interpreted in terms of complex numbers. However, several analytical properties of the complex domain (such as holomorphicity) make the design of CVNNs a more challenging task than their real counterpart. In this paper, we consider the problem of flexible activation functions (AFs) in the complex domain, i.e., AFs endowed with sufficient degrees of freedom to adapt their shape given the training data. While this problem has received considerable attention in the real case, very limited literature exists for CVNNs, where most activation functions are generally developed in a split fashion (i.e., by considering the real and imaginary parts of the activation separately) or with simple phase-amplitude techniques. Leveraging over the recently proposed kernel activation functions, and related advances in the design of complex-valued kernels, we propose the first fully complex, nonparametric activation function for CVNNs, which is based on a kernel expansion with a fixed dictionary that can be implemented efficiently on vectorized hardware. Several experiments on common use cases, including prediction and channel equalization, validate our proposal when compared to real-valued neural networks and CVNNs with fixed activation functions.

Index Terms—Neural networks, activation functions, kernel methods, complex domain.

I. INTRODUCTION

O VER the last years, machine learning techniques have obtained impressive results in a wide range of fields, especially when dealing with supervised problems [1]–[3]. The majority of these applications has focused on the case of real-valued data: as an example, most of the deep learning frameworks currently used today can only work with floating point (or integer) numbers. Several applicable domains of interest, however, exhibit data that can be more naturally modeled using complex-valued algebra, from image processing to time-series prediction, bioinformatics, and robotics’ control (see [4], [5] for a variety of examples). While complex data can immediately be transformed to a real domain by considering the real and imaginary components separately, the resulting loss of phase information gives rise to algorithms that are generally less efficient (or expressive) than alternative methods able to work directly in the complex domain, as evidenced by a large body of literature [6]. Due to this, many learning algorithms have been extended to deal with complex data, including linear adaptive filters [5], [7], kernel methods [8]–[10], component analysis [11], and neural networks (NNs) [12]–[18]. We consider this last class of algorithms here.

Despite the apparent similarity between the real and complex domains, working directly in the latter is challenging because of several non-intuitive analytical properties of the complex algebra. Most notably, almost all cost functions involved in the training of complex models require non-analytic (also known as non-holomorphic) [8] functions, so that standard complex derivatives cannot be used in the definition of the optimization algorithms. This is why several algorithms defined before the last decade considered optimizing the real and imaginary components separately, resulting in a more cumbersome notation which somehow hindered their development [19]. More recently, this problem has been solved by the adoption of the so-called CR-calculus (or Wirtinger’s calculus), allowing to define proper complex derivatives even for non-analytic functions [20], [21], by considering explicitly their dependence on both their arguments and their complex conjugates. We describe CR-calculus more in depth in Section II.

When dealing with neural networks, another challenging task concerns the design of a proper activation function in the complex domain. In the real-valued case, the use of the rectified linear unit (ReLU) has been instrumental in the development of truly deep networks [22], [23], and has spun a wave of further research in the topic, see [24], [25] for very recent examples. In the complex case, Liouville’s theorem asserts that the only complex function which is analytic and bounded at the same time is a constant one. Due to the preference for bounded activation functions before the introduction of the ReLU, many authors in the past preferred bounded functions to analytic ones, most notably in a split organization, wherein the real and imaginary parts of the activations are processed separately [26], or in a phase-amplitude configuration, in which the nonlinearity...
is applied only to the magnitude component, while the phase component is preserved [12]. Even extending the ReLU function to the complex domain has been shown to be non-trivial, and several authors have proposed different variations [14], [16].

In this paper, we consider the problem of adapting activation functions in the complex domain. For real-valued NNs, there is a large body of literature pointing to the fact that endowing activation functions with several degrees of freedom can improve the accuracy of the trained networks, ease the flow of the back-propagated gradient, or vastly simplify the design of the network. In the simplest case, we can consider parametric functions having only a few (generally less than three) parameters per neuron, such as the parametric ReLU [27] or the S-shaped ReLU [28]. More generally, we can think of non-parametric activation functions, that can adapt to potentially any shape in a purely data-driven fashion, with a flexibility that can be controlled by the user, and to which standard regularization techniques can be applied. In the real-valued case, much research has been devoted to the topic, including the design of Maxout networks [29], adaptive piecewise linear (APL) units [30], spline functions [31], and the recently proposed kernel activation functions (KAFs) [32].

When dealing with complex-valued NNs (CVNNs), however, only a handful of works have considered adapting the activation functions, and only in the simplified parametric case [17], or when working in a split configuration [11]. In this sense, how to design activation functions that can adapt to the training data while remaining simple to implement remains an open question.

**A Contributions of the Paper**

We introduce a new family of non-parametric activation functions in the complex domain, building upon the idea of KAFs defined in [32] only in the real domain. In particular, by building on recent works on complex-valued reproducing kernel Hilbert spaces (RKHSs), we propose the first adaptable activation function directly defined in the complex domain. All the functions we introduce can leverage highly vectorized CPU/GPU libraries for matrix multiplication.

The basic idea of KAFs, which were defined in [32] only in the real-valued case, is to exploit a kernel expansion at every neuron, in which the elements of the kernel dictionary are fixed beforehand, while the mixing coefficients are adapted through standard optimization techniques. Here, we propose two different techniques to apply the idea of KAFs in the context of CVNNs. In the first case, we use a split combination where the real and imaginary components are processed by two independent KAFs sharing the same dictionary. In the second case, based on the complex-valued RKHS theory, we are able to redefine the KAF directly in the complex domain, also describing several choices for the kernel function. We show via multiple experimental comparisons that CVNNs endowed with complex-valued KAFs can outperform both real-valued NNs and CVNNs having only fixed or parametric activation functions.

**B. Organization of the Paper**

In Section II we introduce the basic theoretical elements underpinning optimization in a complex domain and CVNNs. Then, in Section III we summarize research on designing activation functions for CVNNs. The two proposed complex KAFs are given in Section IV (split KAF) and Section V (fully complex KAF). We briefly discuss implementation aspects of CVNNs in Section VI. Finally, we provide an experimental evaluation in Section VII before concluding in Section VIII.

**C. Notation**

We denote vectors using boldface lowercase letters, e.g., \( \mathbf{a} \); matrices are denoted by boldface uppercase letters, e.g., \( \mathbf{A} \). All vectors are assumed to be column vectors.

A complex number \( z \in \mathbb{C} \) is represented as \( z = a + ib \), where \( a = \Re\{z\} \) and \( b = \Im\{z\} \) are, respectively, the real part and the imaginary part of the number, and \( i = \sqrt{-1} \). Sometimes, we also use \( z \) and \( z^* \) to denote the real and imaginary parts of \( z \) for simplicity. Magnitude and phase of a complex number are given by \( |z| \) and \( \phi(z) \) respectively. \( z^* = a - ib \) denotes the complex conjugate of \( z \). Other notation is introduced in the text when appropriate.

**II. PRELIMINARIES**

**A. Complex Algebra and CR-Calculus**

We start by introducing the basic theoretical concepts required to define a complex-valued function and to optimize it. We consider scalar functions first, and discuss the multivariate extension later on. Any complex-valued function \( f : \mathbb{C} \rightarrow \mathbb{C} \) can be written as:

\[
    f(z) = u(a, b) + iv(a, b),
\]

where \( u(\cdot, \cdot) \) and \( v(\cdot, \cdot) \) are real-valued functions in two arguments. The function \( f \) is said to be real-differentiable if the partial derivatives of \( u \) and \( v \) with respect to \( a \) and \( b \) are defined. Additionally, the function is called analytic (or holomorphic) if it satisfies the Cauchy-Riemann conditions:

\[
    \frac{\partial u(a, b)}{\partial a} = \frac{\partial v(a, b)}{\partial b} \quad \text{and} \quad \frac{\partial v(a, b)}{\partial a} = -\frac{\partial u(a, b)}{\partial b}.
\]

Only analytic functions admit a complex derivative in the standard sense, but most functions used in practice for CVNNs do not satisfy (2) (such as functions with real-valued outputs for which \( v(a, b) = 0 \) everywhere). In this case, CR-calculus [21] provides a theoretical framework to handle non-analytic functions directly in the complex domain without the need to switch back and forth between definitions in the complex domain and gradients’ computations in the real one.

The main idea of CR-calculus is to consider \( f \) explicitly as a function of both \( z \) and its complex conjugate \( z^* = a - ib \), which we denote as \( f(z, z^*) \). If \( f \) is real-differentiable, then it is also analytic with respect to \( z \) when keeping \( z^* \) constant and vice versa. Thus, we can define a pair of (complex) derivatives as follows [20], [21]:

\[
    \text{R-derivative} \triangleq \left. \frac{\partial f(z, z^*)}{\partial z} \right|_{z^* = \text{const}} = \frac{1}{2} \left( \frac{\partial f}{\partial a} - i \frac{\partial f}{\partial b} \right),
\]

\[
    \text{R*-derivative} \triangleq \left. \frac{\partial f(z, z^*)}{\partial z^*} \right|_{z = \text{const}} = \frac{1}{2} \left( \frac{\partial f}{\partial a} + i \frac{\partial f}{\partial b} \right).
\]
Everything extends to multivariate functions \( f : \mathbb{C}^n \rightarrow \mathbb{C} \) of a complex vector \( \mathbf{z} \in \mathbb{C}^n \) by defining the cogradient and conjugate cogradient operators:

\[
\nabla_{\mathbf{z}} = \left( \frac{\partial}{\partial z_1}, \ldots, \frac{\partial}{\partial z_n} \right)^T, \tag{5}
\]

\[
\nabla_{\mathbf{z}^*} = \left( \frac{\partial}{\partial z_1^*}, \ldots, \frac{\partial}{\partial z_n^*} \right)^T. \tag{6}
\]

Then, a necessary and sufficient condition for \( \mathbf{z}_0 \) to be a minimum of \( f \) is either \( \nabla_{\mathbf{z}_0} f(\mathbf{z}_0, \mathbf{z}_0^*) = 0 \) or \( \nabla_{\mathbf{z}_0} f(\mathbf{z}_0, \mathbf{z}_0^*) = 0 \) [20]. CR-calculus inherits most of the standard properties of the real derivatives, including the chain rule and the differential rule, see [21]. For the important case where the output of the function is real-valued (as is the case for the loss function when optimizing CVNNs) we have the additional property:

\[
\left( \nabla_{\mathbf{z}} f(\mathbf{z}, \mathbf{z}^*) \right)^* = \nabla_{\mathbf{z}} f(\mathbf{z}, \mathbf{z}^*). \tag{7}
\]

Combined with the Taylor expansion of the function, an immediate corollary of this property is that the direction of steepest ascent of \( f \) in the point \( \mathbf{z} \) is given by the conjugate cogradient operator evaluated in that point [21]. Up to a multiplicative constant term, this result coincides with taking the steepest ascent direction with respect to the real derivatives, allowing for a straightforward implementation in most optimization libraries.

### III. Complex-Valued Activation Functions

As we stated in the introduction, choosing a proper activation function in (8) is more challenging than in the real case because of Liouville’s theorem, stating that the only complex-valued functions that are bounded and analytic everywhere are constants. So in practice, one must choose between boundedness and analyticity. Before the introduction of the ReLU activation [22], almost all activation functions in the real case were bounded. Consequently, initial approaches to design CVNNs always preferred non-analytic functions in order to preserve boundedness, most commonly by applying real-valued activation functions separately to the real and imaginary parts [26]:

\[
g(z) = g_R(\Re \{z\}) + ig_R(\Im \{z\}), \tag{12}
\]

where \( z \) is a generic input to the activation function in (8), and \( g_R(\cdot) \) is some real-valued activation function, e.g., sigmoid. This is called a split activation function. As a representative example, the magnitude and phase of the split-tanh when varying the activation are given in Fig. 1. Early proponents of this approach can be found in [36] and [19].

Another common class of non-analytic activation functions are the phase-amplitude (PA) functions popularized by [12], [37]:

\[
g(z) = \frac{z}{c + |z|/r}, \tag{13}
\]

\[
g(z) = \tanh \left\{ \frac{|z|}{m} \right\} \exp \{i\phi(z)\} , \tag{14}
\]

where \( \phi(z) \) is the phase of \( z \), while \( c, r \) and \( m \) are positive constant which in most cases are set equal to 1. PA functions can be seen as the natural generalization of real-valued squashing functions such as the sigmoid, because the output \( g(z) \) has bounded magnitude but preserves the phase of \( z \).

A third alternative is to use fully-complex activation functions that are analytic and bounded almost everywhere, at the cost of introducing a set of singular points. Among all possible transcendental functions, it is common to consider the complex-valued extension of the hyperbolic tangent, defined as [13]:

\[
g(z) = \tanh \{z\} = \frac{\exp \{z\} - \exp \{-z\}}{\exp \{z\} + \exp \{-z\}} , \tag{15}
\]
possessing periodic singular points at the imaginary points $i(0.5 + n)\pi$, with $n \in \mathbb{N}$. However, careful scaling of the inputs and of the initial weights allows to avoid these singularities during training.

Finally, several authors have proposed extensions of the popular real-valued ReLU function $\text{ReLU}(s) = \max\{0, s\}$. As discussed in [17], a simple split configuration as in (12) results in poor performance. An improved complex-valued ReLU is designed in [16] as:

$$g(z) = \begin{cases} z & \text{if } \Re\{z\}, \Im\{z\} \geq 0, \\ 0 & \text{otherwise} \end{cases}.$$  

(16)

Alternatively, inspired by the PA functions to maintain the phase of the activation value, [14] propose the following modReLU function:

$$g(z) = \text{ReLU}(|z| + b) \exp\{i\phi(z)\},$$

(17)

where $b$ is an adaptable parameter defining a radius along which the output of the function is 0. Another extension, the complex cardioid, is advanced in [38]:

$$g(z) = \frac{1}{2} \left(1 + \cos\{\phi(z)\}\right)z,$$

(18)

maintaining phase information while attenuating the magnitude based on the phase itself. For real-valued inputs, (18) reduces to the ReLU.

Note that in all cases these proposed activation functions are fixed or endowed with a very small degree of flexibility (as in (17)). In the following sections we describe a principled technique to design non-parametric activation functions for use in CVNNs.

IV. SPLIT KERNEL ACTIVATION FUNCTIONS

Our first proposal is a split function as in (12), where non-parametric (real-valued) functions for $g_R(\cdot)$ are used in place of fixed ones. Specifically, we consider the kernel activation function (KAF) proposed in [32], which will also serve as a base for the fully complex-valued proposal of the following section. Here, we introduce the basic elements of the KAF, and we refer to the original paper [32] for a fuller exposition.

The basic idea of a KAF is to model each activation function as a one-dimensional kernel model, where the kernel elements are chosen in a proper way to obtain an efficient backpropagation step. Consider the generic activation function $g_R(s)$, where $s$ denotes either the real or the imaginary part of $z$ as in (12). To obtain a flexible shape, we can model a linear predictor on a high-dimensional feature space $\Phi(s)$ of the activation. However, this process becomes infeasible for a large number of feature transformations, and cannot handle infinite-dimensional feature spaces. For feature maps associated to a reproducing kernel Hilbert space $\mathcal{H}$ with kernel $\kappa(\cdot, \cdot)$, we can write an equivalent linear model by exploiting the representer theorem as:

$$g_R(s) = \sum_{n=1}^{D} \alpha_n \kappa(s, d_n),$$

(19)

where $\{\alpha_n\}_{n=1}^{D}$ are the mixing coefficients and $\{d_n\}_{n=1}^{D}$ make up the so-called dictionary of the kernel expansion [39], [40]. Remember that a function $\kappa(\cdot, \cdot)$ is a valid kernel function if it respects the positive semi-definiteness property, i.e., for any possible choice of $\{\alpha_n\}_{n=1}^{D}$ and $\{d_n\}_{n=1}^{D}$ in (19):

$$\sum_{n=1}^{D} \sum_{m=1}^{D} \alpha_n \alpha_m \kappa(d_n, d_m) \geq 0.$$  

(20)

In the context of a neural network, the dictionary elements cannot be selected a priori because they would change at every step of the optimization algorithm depending on the distribution of the activation values. Instead, we exploit the fact that we are working with one-dimensional kernels to fix the elements beforehand, and only adapt the mixing coefficients in the optimization step. In particular, we select the elements $d_1, \ldots, d_D$ by sampling $D$ values over the $z$-axis, uniformly around zero. In this way, the value $D$ becomes a hyper-parameter controlling the flexibility of the approach: for larger $D$ we obtain a more flexible
method at the cost of a larger number of adaptable parameters. In general, since the function is only a small component of a much larger neural network, values in the range \( D \in [10, 20] \) are sufficient for most applications. As the number of parameters per neuron can potentially grow without bound depending on the choice of \( D \), we refer to such activation functions as non-parametric.

The same dictionary is shared across the entire neural network, but with two different sets of mixing coefficients for the real and imaginary parts of each neuron. Due to this, an efficient implementation of the proposed split-KAF is straightforward. In particular, consider the vector \( z \) containing the \( N_l \) (complex) activations of a layer following the linear operations in (8). We build the matrix \( \mathbf{K}_R \in \mathbb{R}^{N_l \times D} \) by computing all the kernel values between the real part of the activations and the elements of the dictionary (and similarly for \( \mathbf{K}_I \) using the imaginary parts), and we compute the final output of the layer as:

\[
\mathbf{h}_l = (\mathbf{A}_R \odot \mathbf{K}_R) \mathbf{1} + i (\mathbf{A}_I \odot \mathbf{K}_I) \mathbf{1},
\]

where \( \odot \) represents element-wise product (Hadamard product), \( \mathbf{A}_R, \mathbf{A}_I \in \mathbb{R}^{N_l \times D} \) are matrices collecting row-wise all the mixing coefficients for the real and imaginary components of the layer, and \( \mathbf{1} \in \mathbb{R}^D \) is a vector of ones. For handling batches of elements (or convolutive layers), we only need to slightly modify (21) by adding additional trailing dimensions.

For all our experiments, we consider the 1D Gaussian kernel defined as:

\[
\kappa(s, d_n) = \exp \left\{ -\gamma (s - d_n)^2 \right\},
\]

where \( \gamma \in \mathbb{R} \) is the inverse of the kernel bandwidth. In the proposed KAF scheme, the values of the dictionary are chosen according to a grid, and as such the optimal bandwidth parameter depends uniquely on the grid resolution. In particular, the following rule-of-thumb was proposed in [32] and it is used in our experiments:

\[
\gamma = \frac{1}{6\Delta^2},
\]

where \( \Delta \) is the distance between the grid points. In order to provide an additional degree of freedom to our method, we also optimize a single \( \gamma \) per layer via back-propagation after initializing it following (23).

V. FULLY-COMPLEX KERNEL ACTIVATION FUNCTIONS

While most of the literature on kernel methods in machine learning has focused on the real-valued case, it is well known that the original mathematical treatment originated in the complex-valued domain [41]. In the context of the kernel filtering literature, techniques to build complex-valued algorithms by separating the real and the imaginary components (as in the previous section) are called complexification methods [8]. However, recently several authors have advocated for the direct use of (pure) complex-valued kernels leveraging the complex-valued treatment of RKHSs for a variety of fields, as surveyed in the introduction.

From a theoretical standpoint, defining complex RKHSs and kernels is relatively straightforward. As an example, a one-dimensional complex-function \( \kappa_C : \mathbb{C} \times \mathbb{C} \to \mathbb{C} \) is positive semi-definite if and only if:

\[
\sum_{n=1}^{D} \sum_{m=1}^{D} \alpha_n \alpha_m \kappa(d_n, d_m) \geq 0, \forall \alpha_n, \alpha_m, d_n, d_m \in \mathbb{C},
\]

where all values are now defined in the complex-domain. Any PSD function is then a valid kernel function. Based on this, in this paper we also propose a fully-complex, non-parametric KAF by defining (19) directly in the complex domain, without the need for split functions:

\[
g(z) = \sum_{n=1}^{D} \sum_{m=1}^{D} \alpha_{n,m} \kappa_C\left(z, d_n + id_m\right),
\]

where the mixing coefficients \( \{\alpha_{n,m}\}_{n,m=1}^{D} \) are now defined as complex numbers. Note that, in order for the dictionary to provide a dense sampling of the space of complex numbers, we now consider \( D^2 \) fixed elements arranged over a regular grid, an example of which is depicted in Fig. 2. Due to this, we now have \( D^2 \) adaptable mixing coefficients per neuron, as opposed to \( 2D \) in the split case. We counter-balance this by selecting a drastically smaller \( D \) (see the experimental section).

An immediate complex-valued extension of the Gaussian kernel in (22) is given by:

\[
\kappa_C(z, d) = \exp \left\{ -\gamma (z - d)^2 \right\},
\]

where in our experiments the bandwidth hyper-parameter \( \gamma \) is selected using the same rule-of-thumb as before and then adapted layer-wise. A complete analysis of the feature space associated to (26) is given in [42]. In order to gain some informal understanding, we can write the kernel explicitly in terms of the
real and imaginary components of its arguments:
\[
\kappa_C(z, d) = \exp \left\{ -\gamma |z_r - d_r|^2 \right\} \exp \left\{ \gamma |z_i + d_i|^2 \right\} \cdot \left( \cos \{ 2\gamma (z_r - d_r)(z_i + d_i) \} - i \sin \{ 2\gamma (z_r - d_r)(z_i + d_i) \} \right).
\]
(27)
By analyzing the previous expression, we see that the complex-valued Gaussian kernel has several properties which are counter-intuitive if one is used to work with its real-valued restriction. First of all, (26) cannot be interpreted as a standard similarity measure, because it depends on its arguments only via \((z_r - d_r)\) and \((z_i + d_i)\). For the same reasons, the kernel is not stationary, and it has an additional oscillatory behavior. We refer to Fig. 3 (or to [10, Section IV-A]) for an illustration of the kernel when fixing the second argument.

For these reasons, another extension of the Gaussian kernel to the complex domain is given in [8], where the authors propose to build a whole family of complex-valued kernels starting from any real-valued one \(\kappa_R\) as follows:
\[
\kappa_C(z, d) = \kappa_R(z_r, d_r) + \kappa_R(z_i, d_i) + i(\kappa_R(z_r, d_i) - \kappa_R(z_i, d_r)).
\]
(28)
The new complex-valued kernel is called an independent kernel. By plugging the real-valued Gaussian kernel (22) in the previous expression, we obtain a complex-valued expression that can still be interpreted as a similarity measure between the two points.

Note that several alternative kernels are also possible, many of which are specific to the complex-valued case, a prominent example being the Szego kernel [8]:
\[
\kappa_C(z, d) = \frac{1}{\left(1 - zd^\ast\right)^2}.
\]
(29)

VI. NOTES ON IMPLEMENTATION
In the previous sections we have described two complete functional models of complex-valued neural networks based on non-parametric activation functions. Nevertheless, the design of the architecture and the training of a CVNN in a practical implementation involve several additional procedures. We now briefly discuss these procedures and comment on how they must be adapted for the complex-valued case w.r.t. real-valued neural networks (RVNNs).

A. Hyperparameter Optimization
The optimization of real-valued hyperparameters (such as hidden layer size) in CVNNs is equivalent to RVNN practices. As such, it can be dealt with by standard hyperparameter optimization methods including grid search, randomized search [43], and Bayesian optimization [44]. The optimization of complex-valued hyperparameters (such a complex step size) has not been explored yet in the literature to the best of our knowledge, and it is beyond the scope of this work. Before considering a fully complex hyperparameter optimization, however, a simple workaround would consist in splitting the complex-valued hyperparameters in real and imaginary part, similar to the strategy followed in Section IV.

B. Weight Initialization

Standard initialization procedures for the linear weights in RVNNs have been described in [45] and [27]. Recently, an extension of these procedures to the complex-valued case was proposed [17], which we adopt in the experiments of Section VII. In particular, we initialize the complex weights of the \(l\)-th layer by drawing their magnitudes from \(N(0, 2N_l)\) and their phases from \(U(-\pi, \pi)\), where \(N_l\) is the number of neurons in this layer.

C. Deep Networks

The construction of deep complex-valued architectures requires overcoming several practical challenges, similar to those that appear in real-valued networks. In the experiments of Section VII, we only consider shallow networks that contain at most three hidden layers. Nevertheless, the interested reader may refer to the discussion on deep complex-valued networks.
in [17], which proposes among others a complex-valued batch normalization technique.

VII. EXPERIMENTAL EVALUATION

In this section, we experimentally evaluate the proposed activation functions on several benchmark problems, including channel identification in Section VII-A, wind prediction in Section VII-B, and multi-class classification in the complex domain in Section VII-C. In all cases, we linearly preprocess the real and imaginary components of the input features to lie in the [−1, +1] range. We regularize all parameters with respect to their squared absolute value (which is equivalent to standard \( \ell_2 \) regularization applied on the real and imaginary components separately), but we exclude the bias terms and the window parameter in (17). We select the strength of the regularization term and the size of the networks based on previous literature or on a cross-validation procedure, as described below. For optimization, we use a simple complex-valued extension of the Adagrad algorithm, which computes a per-parameter learning rate weighted by the squared magnitude of the gradients themselves. For each iteration, we construct a mini-batch by randomly sampling 40 elements from the entire training dataset. All algorithms have been implemented in Python using the Autograd library [46].

A. Experiment 1 - Channel Identification

Our first experiment is a standard benchmark in the complex-valued literature, i.e., a channel identification task [47]. The input to the channel is generated as:

\[
s_n = \left( \sqrt{1 - \rho^2} X_n + i \rho Y_n \right),
\]

where \( X_n \) and \( Y_n \) are Gaussian random variables, and the parameter \( \rho \) determines the circularity\(^1\) of the signal. For \( \rho = \frac{\sqrt{5}}{2} \), the input is circular, while for \( \rho \) approaching 0 or 1 the signal is highly non-circular. The output of the channel is computed by first applying a linear filtering operation:

\[
t_n = \sum_{k=1}^{5} h(k) s_{n-k+1},
\]

where:

\[
h(k) = 0.432 \left( 1 + \cos \left\{ \frac{2\pi(k-3)}{5} \right\} \right) - i \left( 1 + \cos \left\{ \frac{2\pi(k-3)}{10} \right\} \right),
\]

for \( k = 1, \ldots, 5 \). Then, the output of the linear filter goes through a memoryless non-linearity:

\[
r_n = t_n + (0.15 - i0.1) t_n^2,
\]

\(^1\)A random variable \( Z \) is circular if \( Z \) and \( Z \exp\{ i \psi \} \) have the same probability distribution for any angle \( \psi \). Roughly speaking, non-circular signals are harder to predict, requiring the use of widely linear techniques when using standard linear filters [8].

and finally it is corrupted by adding white Gaussian noise in order to get the final signal \( r_n \). The variance of the noise is selected to obtain a signal-to-noise ratio (SNR) of about 13 dB.

The input to the neural network is an embedding of channel inputs:

\[
x = \left[ s_{n-L+1}, s_{n-L+2}, \ldots, s_n \right]^T,
\]

with \( L = 5 \), and the network is trained to output \( r_n \). We generate 2000 samples of the channel, and we randomly keep 15% for testing, averaging over 15 different generations of the dataset. We compare the following algorithms:

- **2R-NN**: a real-valued neural network taking as input the real and imaginary parts separately. For the activation functions in the hidden layers, we consider either a standard \( \tanh \) or ReLUs.
- **C-NN**: complex-valued neural networks with fixed activation functions, including a split-\( \tanh \), a split-ReLU, the AMP function in (13), or the complex ReLU in (16).
- **ModReLU-NN**: CVNN with adaptable activation functions with ModReLU neurons as in (17). In this case, the coefficients of the neurons are all initialized at 0.1 and later adapted.
- **Maxout**: a CVNN where we use the non-parametric Maxout activation function [29] in a split configuration.
- **Proposed KAF-NN**: CVNN with the split-KAF proposed in Section IV. We empirically select \( D = 20 \) elements in the dictionary sampled uniformly in \([-2, +2] \).
- **Proposed C-KAF-NN**: CVNN with the fully complex KAF proposed in Section V. In this case, we test either the complex Gaussian kernel (26), or the independent kernel with the real Gaussian kernel as base. We empirically select \( D = 8 \).

All algorithms are trained by minimizing the mean-squared error in (10) on random mini-batches of 40 elements. Following [34], in this scenario we consider one hidden layer with 10 neurons (as more layers are not found to provide significant improvements in performance). The size of the regularization factor is empirically selected as \( 10^{-4} \). Results in terms of mean squared error (MSE) expressed in dBs are given in Figure 4, by considering either \( \rho = \frac{\sqrt{5}}{2} \) (circular input signal) or the more challenging scenario \( \rho = 0.95 \) (non-circular signal).

As expected, results are generally lower for the non-circular case, proportionally so for techniques that are not able to exploit the geometry of non-circular complex signals, such as non-locally linear models and real-valued neural networks. However, the proposed KAF-NN and C-KAF-NN are able to consistently out-perform all other methods in both scenarios in a stable fashion. Note that this difference in performance cannot be overcome by increasing the size of the other networks, thus pointing to the importance of adapting the activation functions also in the complex case. Interestingly, the complex Gaussian kernel in (26) results in a poor performance, similarly to the split-Maxout, which is solved by using the independent one.
Fig. 4. Results for the first experiment, expressed in terms of MSE (dB). With a dashed line we divide the results of the proposed models.

Fig. 5. A plot of the complex-valued wind profile for the initial 500 samples of the wind time-series.

### TABLE I

<table>
<thead>
<tr>
<th>Model</th>
<th>$R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Linear</td>
<td>0.361 ± 0.0227</td>
</tr>
<tr>
<td>2R-NN (tanh)</td>
<td>0.424 ± 0.015</td>
</tr>
<tr>
<td>2R-NN (ReLU)</td>
<td>0.435 ± 0.016</td>
</tr>
<tr>
<td>C-NN (split-tanh)</td>
<td>0.426 ± 0.015</td>
</tr>
<tr>
<td>C-NN (split-ReLU)</td>
<td>0.438 ± 0.016</td>
</tr>
<tr>
<td>C-NN (AMP)</td>
<td>0.431 ± 0.014</td>
</tr>
<tr>
<td>C-NN (CRELU)</td>
<td>0.181 ± 0.106</td>
</tr>
<tr>
<td>ModReLU-NN</td>
<td>0.438 ± 0.015</td>
</tr>
<tr>
<td>KAF-NN</td>
<td>0.444 ± 0.015</td>
</tr>
<tr>
<td>C-KAF-NN</td>
<td>0.424 ± 0.016</td>
</tr>
<tr>
<td>C-KAF-NN (Ind.)</td>
<td>0.442 ± 0.016</td>
</tr>
</tbody>
</table>

B. Experiment 2 - Wind Prediction

For the second experiment, we consider a real-world dataset for a task of wind prediction [48]. The dataset consists of 5000 hourly samples of wind intensity collected along two different axes (north axis and east axis). The dataset is provided in three settings of wind regime, namely ‘low’, ‘medium’, and ‘high’, from which we select the highest, being the most challenging one. In order to construct a complex-valued signal, the two samples for each hour are considered as the real and the imaginary components of a single complex number (for more motivation on the use of complex-valued information when dealing with wind forecasting, see [18], [48]–[51]). A snapshot of the absolute value and phase of the resulting signal is shown in Fig. 5 for the initial 500 samples. We consider the task of predicting both components of the wind for an 8-hour-ahead horizon, starting from an embedding of the last 10 hours of measurements. We select neural networks with 2 hidden layers (as more hidden layers are not found to provide gain in performance), and we optimize both the number of neurons and the regularization factor on a held-out validation set. We test the datasets on the last 500 components of the time-series, in terms of the $R^2$ coefficient of determination:

$$R^2 = 1 - \frac{\sum_{n=1}^{500} |y_n - \hat{y}_n|^2}{\sum_{n=1}^{500} |y_n - \bar{y}|^2},$$  \hspace{1cm} (35)

where $y_n$ is the true value, $\hat{y}_n$ is the predicted value, and $\bar{y}$ is the mean of the true values computed from the test set. Positive
values of $R^2$ denotes a prediction which is better than chance, with values approaching 1 for an almost-perfect prediction.

Results for the experiment are reported in Table I. We can see that, also in this scenario, the two best results are obtained by the proposed split-KAF and complex KAF neurons, significantly outperforming the other models.

### C. Experiment 3: Complex-Valued Multi-Class Classification

We conclude our experimental evaluation by testing the proposed algorithms on four multi-class image classification problems expressed in the complex domain. Following [47], we build each task by applying a two-dimensional fast Fourier transform (FFT) to the images in the well-known MNIST dataset, comprising 60000 28 × 28 black-and-white images of handwritten digits split into ten classes. We then rank the coefficients of the FFT in terms of significance (by considering their mean absolute value), and keep only the 100 most significant coefficients as input to the models. In order to provide a wider comparison, we also apply the same procedure to three additional datasets:

- **Fashion MNIST** (F-MNIST) [52]: a variant of MNIST concerning images of clothing items, with the same dimensionality.
- **Extended MNIST** (EMNIST) [53]: a set of extensions of MNIST, from which we consider the ‘Digits’ one, comprising 240 thousand images of handwritten digits.
- **Latin OCR** [54]: an OCR problem concerning handwritten Latin digits segmented from real manuscripts of the Vatican secret archives. There are approximately 12 thousands characters belonging to 23 separate classes.

We compare a real-valued NN taking the real and the imaginary components of the coefficients as separate inputs, a CVNN with modReLU activation functions, and two CVNNs employing split-KAFs and fully-complex KAFs with independent kernels. All networks have a softmax activation function in their output layer. For the CVNNs, we use the following variation to handle the complex valued activations $h$:

$$ \text{softmax}_n(h) = \frac{\exp \left\{ \Re \{ h_n \}^2 + \Im \{ h_n \}^2 \right\}}{\sum_{i=1}^{C} \exp \left\{ \Re \{ h_i \}^2 + \Im \{ h_i \}^2 \right\}}, $$

where $h \in \mathbb{C}^C$, and $C = 10$ for our problem. All networks are then trained by minimizing the classical regularized cross-entropy formulation with the same optimizer as the last sections. We consider networks with three hidden layers having 100 neurons each, whose regularization term is optimized via cross-validation separately. We also apply an early stopping procedure (with the standard splits from each dataset), stopping whenever accuracy is not improving for 1000 iterations of optimization. Results on the test sets are provided in Table II.

We see that working in the complex domain results in significantly better performance when compared to working in the real domain. We show a representative evolution of the loss function for MNIST in Fig. 6, where we highlight the first 10000 iterations for readability.

### VIII. CONCLUSIVE REMARKS

In this paper, we considered the problem of adapting activation functions in a complex-valued neural network (CVNN). To this end, we proposed two different non-parametric models that extend the recently introduced kernel activation function (KAF) to the complex-valued case. The first model is a split configuration, where the real and the imaginary components of the activation are processed independently by two separate KAFs. In the second model, we directly redefine the KAF in
the complex domain with the use of fully-complex kernels. We showed that CVNNs with adaptable functions can outperform neural networks with fixed functions in different benchmark problems, including channel identification, wind prediction, and multi-class classification. For the fully-complex KAF, the independent kernel generally outperforms a naive complex Gaussian kernel without introducing significantly more complexity.

Due to the space constraints, in this paper we have focused only on a selected number of experimental comparisons, with a limited number of complex kernels. In order to overcome these limitations, multiple future works are possible, most notably by leveraging over recent advances in the field of real-valued kernels [55] and complex-valued kernel regression and classification. One example is the use of pseudo-kernels [10] to handle more efficiently the non-circularity in the signals propagated through the network. Additionally, we plan on comparing with more datasets, e.g., [10], and analyze the temporal convergence rate comparison of CVNN with real-valued approaches. More generally, it would be interesting to extend other classes of non-parametric, real-valued activation functions (such as Maxout networks [29] or adaptive piecewise linear units [30]) to the complex domain, or adapt the proposed complex KAFs to other types of NNs, such as convolutive architectures [1], [56].

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REFERENCES


Simone Scardapane received the B.Sc. degree in computer engineering from Roma Tre University, Rome, Italy in 2009, the M.Sc. degree in artificial intelligence and robotics in 2011, and the Ph.D. degree from Sapienza University of Rome, Rome, Italy in 2016, researching mainly in the fields of distributed machine learning and adaptive audio processing. Before receiving the Ph.D. degree, he was a Software/Web Developer for one year. He is currently a Postdoctoral Fellow with Sapienza University, and an honorary Research Fellow at the CogBID laboratory with the University of Stirling, Stirling, U.K.

Steven Van Vaerenbergh (M’11–SM’15) received the M.Sc. degree in electrical engineering from Ghent University, Ghent, Belgium in 2003, and the Ph.D. degree from the University of Cantabria, Santander, Spain in 2010. He was a Visiting Researcher with the Computational Neuroengineering Laboratory, University of Florida, Gainesville, FL, USA in 2008. He is currently a Postdoctoral Associate with the Department of Telecommunications Engineering, University of Cantabria, Santander, Spain. His research interests include machine learning algorithms for pattern recognition, prediction, system identification, and online machine learning.

Amir Hussain received the B.Eng. and Ph.D. degrees from the University of Strathclyde, Glasgow, U.K., in 1992 and 1997, respectively.

He is currently a Professor and the founding Head of the Cyber and Big Data Research Laboratory, Edinburgh Napier University, Edinburgh, U.K. He has coauthored more than 320 papers, including more than a dozen books and around 120 journal papers. He is a founding Editor-in-Chief for the journals Cognitive Computation (Springer Nature) and Big Data Analytics (BioMed Central), and of the Springer book series on Socio-Affective Computing and Cognitive Computation Trends. He is currently an Associate Editor for the IEEE TRANSACTIONS ON NEURAL NETWORKS AND LEARNING SYSTEMS, the IEEE COMPUTATIONAL INTELLIGENCE MAGAZINE, and the IEEE TRANSACTIONS ON EMERGING TOPICS IN COMPUTATIONAL INTELLIGENCE. Among other distinguished roles, he is the General Chair for IEEE WCCI 2020, and the Vice-Chair of the Emergent Technologies Technical Committee of the IEEE Computational Intelligence Society.

Aurelio Uncini (M’88) received the Laurea degree in electronic engineering from the University of Ancona, Ancona, Italy, in 1983, and the Ph.D. degree in electrical engineering from the University of Bologna, Bologna, Italy, in 1994. He is currently a Full Professor with the Department of Information Engineering, Electronics and Telecommunications, where he is teaching neural networks, adaptive algorithm for signal processing, and digital audio processing, and where he is the Founder and the Director of the “Intelligent Signal Processing and Multimedia” group. He is a member of the Associazione Elettrotecnica ed Elettronica Italiana, of the International Neural Networks Society, and of the Societ Italiana Reti NeuroniChe.