NONLINEAR SIGNAL PROCESSING BASED ON REPRODUCING KERNEL HILBERT SPACE

By

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To my parents, friends and teachers
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My research aimed at analyzing the recently proposed correntropy function and presents a new centered correntropy function from time-domain and frequency-domain approaches. It demonstrates that correntropy and centered correntropy functions not only capture the time and space structures of signals, but also partially characterize the higher order statistical information and nonlinearity intrinsic to random processes. Correntropy and centered correntropy functions have rich geometrical structures. Correntropy is positive definite and centered correntropy is non-negative definite, hence by Moore-Aronszajn theorem they uniquely induce reproducing kernel Hilbert spaces. Correntropy and centered correntropy functions combine the data dependent expectation operator and data independent kernels to form another data dependent operator. Correntropy and centered correntropy functions can be formulated as “generalized” correlation and covariance functions on nonlinearily transformed random signals via the data independent kernel functions. Those nonlinearily transformed signals appear on the sphere in the reproducing kernel Hilbert space induced by the kernel functions if isotropic kernel functions are used. The other approach is to directly work with the reproducing kernel Hilbert space induced by the correntropy and centered correntropy functions directly. The nonlinearily transformed signals in the reproducing kernel Hilbert space is no longer stochastic but rather deterministic. The reproducing kernel Hilbert space induced by the correntropy and centered correntropy functions includes the expectation
operator as embedded vectors. The two views further our understandings of correntropy and centered correntropy functions in geometrical perspective. The two reproducing kernel Hilbert space induced by kernel functions and correntropy functions respectively represent stochastic and deterministic functional analysis.

The correntropy dependence measure is proposed based on the correntropy coefficient as a novel statistical dependence measure. The new measure satisfies all the fundamental desirable properties postulated by Renyi. We apply the correntropy concept in pitch determination, and nonlinear component analysis. The correntropy coefficient is also employed as a novel similarity measure to quantify the inder-dependencies of multi-channel signals.
CHAPTER 1
INTRODUCTION

1.1 Definition of Reproducing Kernel Hilbert Space (RKHS)

A reproducing kernel Hilbert space (RKHS) is a special Hilbert space associated with a kernel such that reproduces (via an inner product) each function in the space, or, equivalently, every point evaluation functional is bounded. Let $\mathcal{H}$ be a Hilbert space of functions on some set $E$, define an inner product $\langle \cdot, \cdot \rangle_{\mathcal{H}}$ in $\mathcal{H}$ and a complex-valued bivariate function $\kappa(x, y)$ on $E \times E$. Then the function $\kappa(x, y)$ is said to be positive definite if for any finite point set $\{x_1, x_2, \ldots, x_n\} \in E$ and for any not all zero corresponding complex number $\{\alpha_1, \alpha_2, \ldots, \alpha_n\} \in \mathbb{C}$,

$$\sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \overline{\alpha_j} \kappa(x_i, x_j) > 0. \quad (1-1)$$

Any positive definite bivariate function $\kappa(x, y)$ is a reproducing kernel because of the following fundamental theorem.

**Moore-Aronszajn Theorem**: Given any positive definite function $\kappa(x, y)$, there exists a uniquely determined (possibly finite dimensional) Hilbert space $\mathcal{H}$ consisting of functions on $E$ such that

(i) for every $x \in E$, $\kappa(x, \cdot) \in \mathcal{H}$ and

$$\sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \overline{\alpha_j} \kappa(x_i, x_j) > 0. \quad (1-2)$$

(ii) for every $x \in E$ and $f \in \mathcal{H}$, $f(x) = \langle f, \kappa(x, \cdot) \rangle_{\mathcal{H}}. \quad (1-3)$

Then $\mathcal{H} := \mathcal{H}(\kappa)$ is said to be a reproducing kernel Hilbert space with reproducing kernel $\kappa$. The properties (1–2) and (1–3) are called the *reproducing property* of $\kappa(x, y)$ in $\mathcal{H}(\kappa)$.

The reproducing kernel Hilbert space terminology has existed for a long time since all the Green’s functions of self-adjoint ordinary differential equations and some bounded Green’s functions in partial differential equations belong to this type. But it is not until 1943 that N. Aronszajn [1] systematically developed the general theory of RKHS and named the term “reproducing kernel”. The expanded paper [2] on his previous work
became the standard reference for RKHS theory. Around the same time, there are some independent work on RKHS presented in the Soviet Union. For instances, A. Povzner derived many of the basic properties of RKHS in [3] and presented some examples in [4]. Meanwhile M.G. Krein also derived some RKHS properties in his study of kernels with certain invariance conditions [5]. Other works studying RKHS theory include Hille [6], Meschkowski [7], Shapiro [8], Saitoh [9] and Davis [10]. Bergman introduced reproducing kernels in one and several variables for the classes of harmonic and analytic functions [11]. He applied the kernel functions in the theory of functions of one and several complex variables, in conformal mappings, pseudo-conformal mappings, invariant Riemannian metrics and other subjects. A more abstract development of RKHS appears in a paper by Schwarts [12].

As discussed in the Moore-Aronszajn Theorem, RKHS theory and the theory of positive definite functions are two sides of the same coin. In 1909, J. Mercer examined the positive definite functions satisfying in Eq. (1–1) in the theory of integral equations developed by Hilbert. Mercer proved that positive definite kernels have nice properties among all the continuous kernels of integral equations [13]. This was the celebrated Mercer’s Theorem which became the theoretic foundation of application of RKHS in statistical signal processing and machine learning. E. H. Moore also studied those kernels in his general analysis context under the name of positive Hermitian matrix and discovered the fundamental theorem above [14]. Meanwhile, S. Bochner examined continuous functions \( \phi(x) \) of a real variable \( x \) such that \( \kappa(x, y) = \phi(x - y) \) satisfying condition in his studying of Fourier transformation. He named such functions positive definite functions [15]. Later A. Weil, I. Gelfand, D. Raikov and R. Godement generalized the notion in their investigations of topological groups. These functions were also applied to the general metric geometry by I. Schoenberg [16, 17], J. V. Neumann and others. In [18], J. Stewart provides a concise historical survey of positive definite functions and their principal generalizations as well as a useful bibliography.
From a mathematical perspective, the one-to-one correspondence between the RKHS and positive definite functions relates operator theory and the theory of functions. It finds useful applications in numerous fields, of which includes: orthogonal polynomials, Gaussian processes, harmonic analysis on semigroups, approximation in RKHS, inverse problem, interpolation, zero counting polynomials, and etc. The book [19] offers a review of recent advance in RKHS in many mathematical fields.

More relevant to this proposal is the RKHS methods in probability theory, random processes and statistical learning theory. I will present a brief review on these two in the following sections separately.

1.2 RKHS in Statistical Signal Processing

Almost all the literature dealing with RKHS methods in statistical signal processing only considered the second order random processes. The rational behind this is that random processes can be approached by purely geometric methods when they are studied in terms of their second order moments - variances and covariances [20].

Given a probability space \((\Omega, \mathcal{F}, P)\), we can define a linear space \(L^2(\Omega, \mathcal{F}, P)\) to be the set consisting all the random variables \(X\) whose second moment satisfying

\[
E[|X|^2] = \int_{\Omega} |X|^2 dP < \infty.
\]  

(1-4)

Furthermore we can impose an inner product between any two random variables \(X\) and \(Y\) in \(L^2(\Omega, \mathcal{F}, P)\) as

\[
\langle X, Y \rangle = E[XY] = \int_{\Omega} XY dP.
\]  

(1-5)

Then \(L^2(\Omega, \mathcal{F}, P)\) becomes an inner product space. Moreover it possesses the completeness property in the sense that the Cauchy sequence converges in the space itself [21]. Hence the inner product space \(L^2(\Omega, \mathcal{F}, P)\) of all square integrable random variables on the probability space \((\Omega, \mathcal{F}, P)\) is a Hilbert space.
Consider a second order random process \( \{ x_t : t \in T \} \) defined on a probability space \( (\Omega, \mathcal{F}, P) \) satisfying
\[
E [ | x_t |^2 ] = \int_{\Omega} | x_t |^2 \, dP < \infty \quad \forall \, t \in T
\] (1–6)
for the second moment of all the random variables \( x_t \). Then for each \( t \in T \), the random variable \( x_t \) can be regarded as a data point in the Hilbert space \( L_2(\Omega, \mathcal{F}, P) \). Hilbert space can thus be used to study the random processes. In particular, we can consider constructing a Hilbert space spanned by a random process.

We define a linear manifold for a given random process \( \{ x_t : t \in T \} \) to be the set of all random variables \( X \) which can be written in the form of linear combinations
\[
X = \sum_{k=1}^{n} c_k x_{t_k}
\] (1–7)
for any \( n \in \mathbb{N} \) and \( c_k \in \mathbb{C} \). Close the set in Eq. (1–7) topologically according to the convergence in the mean using the norm
\[
\| Y - Z \| = \sqrt{E [ | Y - Z |^2 ]}
\] (1–8)
and denote the set of all linear combinations of random variables and its limit points by \( L_2(x_t, t \in T) \). By the theory of quadratically integrable functions, we know that the linear space \( L_2(x_t, t \in T) \) forms a Hilbert space if an inner product is imposed by the definition of Eq. (1–5) with corresponding norm of Eq. (1–8). Notice that \( L_2(x_t, t \in T) \) is included in the Hilbert space of all quadratically integrable functions on \( (\Omega, \mathcal{F}, P) \), hence
\[
L_2(x_t, t \in T) \subseteq L_2(\Omega, \mathcal{F}, P).
\]
Indeed, it can be a proper subset. Therefore by studying the Hilbert space \( L_2(x_t, t \in T) \) we can gain the knowledge of the Hilbert space \( L_2(\Omega, \mathcal{F}, P) \). One of the theoretic foundations to employ RKHS approach to study second order random processes is that the covariance function of random processes induces a reproducing kernel Hilbert space and there exists an isometric isomorphism (congruence) between \( L_2(x_t, t \in T) \) and
the RKHS determined by its covariance function. It was Kolmogorov who first used Hilbert space theory to study random processes [22]. But it was until in the late 1940s that Loève established the first link between random processes and reproducing kernels [23]. He pointed out that the covariance function of a second-order random process is a reproducing kernel and vice versa. Loève also presented the basic congruence (isometric isomorphism) relationship between the RKHS induced by the covariance function of a random process and the Hilbert space of linear combinations of random variables spanned by the random process [24].

Consider two abstract Hilbert space \( \mathcal{H}_1 \) and \( \mathcal{H}_2 \) with inner products denoted as \( \langle f_1, f_2 \rangle_1 \) and \( \langle g_1, g_2 \rangle_2 \) respectively, \( \mathcal{H}_1 \) and \( \mathcal{H}_2 \) are said to be isomorphic if there exists a one-to-one and surjective mapping \( \psi \) from \( \mathcal{H}_1 \) to \( \mathcal{H}_2 \) satisfying the following properties

\[
\psi(f_1 + f_2) = \psi(f_1) + \psi(f_2) \quad \text{and} \quad \psi(\alpha f) = \alpha \psi(f)
\] (1–9)

for all functionals in \( \mathcal{H}_1 \) and any real number \( \alpha \). The mapping \( \psi \) is called an isomorphism between \( \mathcal{H}_1 \) and \( \mathcal{H}_2 \). The Hilbert spaces \( \mathcal{H}_1 \) and \( \mathcal{H}_2 \) are said to be isometric if there exists a mapping \( \psi \) that preserves inner products,

\[
\langle f_1, f_2 \rangle_1 = \langle \psi(f_1), \psi(f_2) \rangle_2,
\] (1–10)

for all functions in \( \mathcal{H}_1 \). A mapping \( \psi \) satisfying both properties Eq. (1–9) and Eq. (1–10) is said to be an isometric isomorphism or congruence. The congruence maps both linear combinations of functionals and limit points from \( \mathcal{H}_1 \) into corresponding linear combinations of functionals and limit points in \( \mathcal{H}_2 \) [20].

Given a second-order random process \( \{x_t : t \in \mathbb{T}\} \) satisfying Eq. (1–6), we know that the mean value function \( \mu(t) \) is well defined according to the Cauchy-Schwartz inequality. We can always assume that \( \mu(\cdot) \equiv 0 \), if not we can preprocess the random process to reduce the DC component. The covariance function is defined as

\[
R(t, s) = E[x_t x_s]
\] (1–11)
which is also equal to the auto-correlation function. It is well known that the covariance
function $R$ is non-negative definite, therefore it determines a unique RKHS, $\mathcal{H}(R)$,
according to the Moore-Aronszajn Theorem. We can construct the RKHS induced by
the covariance function $R$ in the following procedure. First, a series expansion to the
covariance function $R$ can be found by the Mercer’s theorem.

**Mercer’s Theorem:** Suppose $R(t, s)$ is a continuous symmetric non-negative
function on a closed finite interval $\mathbb{T} \times \mathbb{T}$. Denote by $\{\lambda_k, k = 1, 2, \ldots\}$ a sequence
of non-negative eigenvalues of $R(t, s)$ and by $\{\varphi_k(t), k = 1, 2, \ldots\}$ the sequence of
corresponding normalized eigenfunctions, in other word, for all integers $t$ and $s$,

\[
\int \mathbb{T} R(t, s)\varphi_k(t)dt = \lambda_k\varphi_k(s), \quad s, t \in \mathbb{T}
\]  

(1–12)

\[
\int \mathbb{T} \varphi_k(t)\varphi_j(t)dx = \delta_{k,j}
\]  

(1–13)

where $\delta_{k,j}$ is the Kronecker delta function, i.e., equal to 1 or 0 according as $k = j$ or $k \neq j$.
Then

\[
R(t, s) = \sum_{k=0}^{\infty} \lambda_k \varphi_k(t)\varphi_k(s)
\]  

(1–14)

where the series above converges absolutely and uniformly on $\mathbb{T} \times \mathbb{T}$ [13].

Then we can define a function $f$ on $\mathbb{T}$ as the form of

\[
f(t) = \sum_{k=0}^{\infty} \lambda_k a_k \varphi_k(t),
\]  

(1–15)

where the sequence $\{a_k, k = 1, 2, \ldots\}$ satisfies the following condition

\[
\sum_{k=0}^{\infty} \lambda_k a_k^2 < \infty.
\]  

(1–16)

Let $\mathcal{H}(R)$ be the set composed of functions $f(\cdot)$ which can be represented in the form Eq.
(1–15) in terms of eigenfunctions $\varphi_k$ and eigenvalues $\lambda_k$ of the covariance function $R(t, s)$.  

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Furthermore we might define an inner product of two functions in $\mathcal{H}(R)$ as

$$\langle f, g \rangle = \sum_{k=0}^{\infty} \lambda_k a_k b_k,$$  \hspace{1cm} (1–17)  

where $f$ and $g$ are of form Eq. (1–15) and $a_k, b_k$ satisfy property Eq. (1–16). One might as well show $\mathcal{H}(R)$ is complete. Let $f_n(t) = \sum_{k=0}^{\infty} \lambda_k a_k^{(n)} \varphi_k(t)$ be a Cauchy sequence in $\mathcal{H}(R)$ such that each sequence $\{a_k^{(n)}, n = 1, 2, \ldots\}$ converges to a limit point $a_k$. Hence the Cauchy sequence converges to $f(t) = \sum_{k=0}^{\infty} \lambda_k a_k \varphi_k(t)$ which belongs to $\mathcal{H}(R)$. Therefore $\mathcal{H}(R)$ is a Hilbert space. $\mathcal{H}(R)$ has two important properties which make it a reproducing kernel Hilbert space. First, let $R(t, \cdot)$ be the function on $\mathbb{T}$ with value at $s$ in $\mathbb{T}$ equal to $R(t, s)$, then by the Mercer’s Theorem eigen-expansion for the covariance function Eq. (1–14), we have

$$R(t, s) = \sum_{k=0}^{\infty} \lambda_k a_k \varphi_k(s), \quad a_k = \varphi_k(t).$$  \hspace{1cm} (1–18)  

Therefore, $R(t, \cdot) \in \mathcal{H}(R)$ for each $t$ in $\mathbb{T}$. Second, for every function $f(\cdot) \in \mathcal{H}(R)$ of form given by Eq. (1–15) and every $t$ in $\mathbb{T}$,

$$\langle f, R(t, \cdot) \rangle = \sum_{k=0}^{\infty} \lambda_k a_k \varphi_k(t) = f(t).$$  \hspace{1cm} (1–19)  

By the Moore-Aronszajn Theorem, $\mathcal{H}(R)$ is a reproducing kernel Hilbert space with $R(t, s)$ as the reproducing kernel. It follows that

$$\langle R(t, \cdot), R(s, \cdot) \rangle = \sum_{k=0}^{\infty} \lambda_k \varphi_k(t) \varphi_k(s) = R(t, s).$$  \hspace{1cm} (1–20)  

Thus $\mathcal{H}(R)$ is a representation of the random process $\{x_t : t \in \mathbb{T}\}$ with covariance function $R(t, s)$. One may define a congruence $\mathcal{G}$ from $\mathcal{H}(R)$ onto $L_2(x_t, t \in \mathbb{T})$ such that

$$\mathcal{G}(R(t, \cdot)) = x_t.$$  \hspace{1cm} (1–21)
In order to obtain an explicit representation of $\mathcal{G}$, we define an orthogonal random variable sequence $\{\xi_m, m = 1, 2, \ldots\}$ such that

$$E[\xi_k \xi_m] = \begin{cases} 0, & k \neq m \\ \lambda_k, & k = m \end{cases},$$

where $\lambda_k$ and $\psi_k(f_i)$ are eigenvalue and eigenfunction associated with the kernel function $R(t, s)$ by the Mercer’s theorem. We achieve an orthogonal decomposition of the random process as

$$x_t = \sum_{k=0}^{\infty} \varphi_k(t) \xi_k, \quad \forall f(x) \in \mathcal{E}. \quad (1–22)$$

Note that the congruence map $\mathcal{G}$ can be characterized as the unique mapping from $\mathcal{H}(R)$ onto $L_2(x_t, t \in \mathbb{T})$ satisfying the condition that for every functional $f$ in $\mathcal{H}(R)$

$$E[\mathcal{G}(f)x_t] = \langle f, R(t, \cdot) \rangle = f(t). \quad (1–23)$$

It is obvious that $\mathcal{G}$ in Eq. (1–21) fulfills the condition Eq. (1–23). Then the congruence map can be represented explicitly as

$$\mathcal{G}(f) = \sum_{k=0}^{\infty} a_k \xi_k, \quad \forall f \in \mathcal{H}(R), \quad (1–24)$$

where $a_k$ satisfies condition Eq. (1–16).

To prove the representation Eq. (1–24) is a valid and unique map, substituting Eq. (1–22) and Eq. (1–24) into Eq. (1–23), we obtain

$$E[\mathcal{G}(f)x_t] = E \left[ \sum_{k=0}^{\infty} a_k \xi_k \sum_{m=0}^{\infty} \varphi_m(t) \xi_m \right] = \sum_{k=0}^{\infty} \sum_{m=0}^{\infty} a_k \varphi_m(t) E[\xi_k \xi_m]$$

$$= \sum_{k=0}^{\infty} \lambda_k a_k \varphi_k(t) = f(t). \quad (1–25)$$

Parzen applied Loève’s results to statistical signal processing, particularly the estimation, regression and detection problems in late 1950s. Parzen clearly illustrated that the RKHS approach offers an elegant general framework for minimum variance unbiased
estimation of regression coefficients, least-squares estimation of random variables, and
detection of know signals in Gaussian noise. Actually, the solutions to all these problems
can be written in terms of RKHS inner product. Parzen [25] derived the basic RKHS
formula for the likelihood ratio in detection through sampling representations of the
observed random process. The nonsingularity condition for the known signal problem was
also presented. In [26], Parzen provided a survey on the wide range of RKHS applications
in statistical signal processing and random processes theory. The structural equivalences
among problems in control, estimation, and approximation are briefly discussed. These
research directions have been developed further since 1970. Most of Parzen’s results can be

Meanwhile, a Czechoslovakia statistician named Hájek established the basic
congruence relation between the Hilbert space of random variables spanned by a random
process and the RKHS determined by the covariance function of the random process
unaware of the work of Loève, Parzen and even Aronszajn. In a remarkable paper [30],
he shows that the estimation and detection problems can be approached by inverting the
basic congruence map for stationary random processes with rational spectral densities.
Hájek also derived the likelihood ratio using only the individual RKHS norms under a
strong nonsingularity condition.

In early 1970s, Kailath presented a series of papers on RKHS approach to detection
and estimation problems [31–35]. In paper [31], Kailath discusses the RKHS approach
in great details to demonstrate its superiority in computing likelihood ratios, testing for
nonsingularity, bounding signal detectability, and determining detection stability. A simple
but formal expression for the likelihood ratio using RKHS norm is presented in paper [32].
It also presents a test that can verify the likelihood ratio obtained from the formal RKHS
expressions is correct. The RKHS approach to detection problems is based on the fact
that the statistics of a zero-mean Gaussian random process are completely characterized
by its covariance function, which turns out to be a reproducing kernel. In order to extend
to Non-Gaussian random processes detection, characteristic function is used to represent the Non-Gaussian process since it completely specifies the statistics and it is symmetric, non-negative definite and thus a reproducing kernel. Duttweiler and Kailath generalize the RKHS work to Non-Gaussian processes in [34]. Paper [35] considers the variance bounds for unbiased estimates of parameters determining the mean or covariance of a Gaussian random process. An explicit formula is also provided for estimating the arrival time of a step function in white Gaussian noise.

RKHS method is also applied to more difficult aspects of random processes. For instances, Hida and Ikeda study the congruence relation between the nonlinear span of an independent increment process and the RKHS determined by its characteristic function. Orthogonal expansions of nonlinear functions of such processes can be derived based on this relation [36]. Kallianpur presents a nonlinear span expression for a Gaussian process as the direct sum of tensor product [37]. Another important RKHS application area is the canonical, or innovations, representations for Gaussian processes. Hida was the first to present the connection of RKHS and canonical representations [38]. Kailath presented the generalized innovations representations for non-white noise innovations process [33]. RKHS has also been applied to deal with Markovian properties of multidimensional Gaussian processes (random fields). Paper [39, 40] provide RKHS development on multidimensional Brownian motion and the conditions for more general Gaussian fields to be Markovian.

Besides the successful applications of RKHS in estimation, detection and other statistical signal processing areas, there have been extensive research on applying RKHS to a wide variety of problems in optimal approximation including interpolation and smoothing by spline functions in one or more dimensions (curve and surface fitting). In [41] Weinert surveys the one-dimensional case in RKHS formulation of recursive spline algorithms and connections with least-square estimation. Optimality properties of splines are developed and an explicit expression for the reproducing kernel in the polynomial case is proposed by de Boor in [42]. Schumaker presents a survey of applications of RKHS
in multidimensional spline fitting in [43]. Wahba presents extensive results on spline in [44]. Figueiredo took a different approach to apply RKHS in nonlinear system and signal analysis. He built the RKHS from bottom-up using arbitrarily weighted Fock spaces [45]. The spaces are composed of polynomials or power series in either scalar variable or multi-dimensional ones. The spaces can also be extended to infinite or finite Volterra functional or operator series. The generalized Fock spaces have been applied to nonlinear system approximation, semiconductor device characteristics modeling and others [45].

The RKHS approach has enjoyed its successful applications in a wide range of problems in statistical signal processing since 1940s, and continues bringing new perspectives and methods towards old and new problems. The essential idea behind this is that there exits a congruence map between the Hilbert space of random variables spanned by the random process and its covariance function which determines a unique RKHS. The RKHS framework provides a natural link between stochastic and deterministic functional analysis.

1.3 RKHS in Statistical Learning Theory

The statistical learning theory is the mathematical foundation for a broad range of learning problems including pattern recognition, regression estimation, density estimation and etc. The general definition of a learning problem can be stated as follows. Given a set of independent identically distributed (i.i.d.) random variable $x$ drawn from a fixed but unknown distribution $P(x)$, a corresponding set of output random variable $y$ for every input $x$ according to a fixed but unknown conditional distribution $P(y|x)$, and a learning machine that can implement a set of functions $f(x, \lambda)$, $\lambda \in \Lambda$, the problem of learning from examples is to select the function $f(x, \lambda)$ to predict the output response in the best possible way. One employs the loss or discrepancy measure $L(y, f(x, \lambda))$ between the output $y$ given the input $x$ and the response of $f(x, \lambda)$ from the learning machine to select the best function. In the statistical learning theory, mainly developed by Vapnik and Chervonenkis in 1990s [46, 47], the risk minimization criterion is used to search for the
best function. The risk functional which characterizes the loss measure is given by

$$T(\lambda) = \int L(y, f(x, \lambda))dP(x, y).$$

(1–26)

The objective is to find the optimal function $f(x, \lambda_o)$ such that the risk functional $R(\lambda)$ is minimized over all the possible functions when the joint probability distribution $P(x, y)$ is fixed but unknown and the only available information is the data set.

The evolution of statistical learning theory has undergone three periods. In the 1960s efficient linear algorithms were proposed to detect linear relations between the input and response. One example was the perceptron algorithm which was introduced in 1958 [48]. The major research challenge at that time was the problem of how to detect the nonlinear relations. In the mid 1980s, the field of statistical learning underwent a nonlinear revolution with the almost simultaneous introduction of backpropagation multilayered neural networks and efficient decision tree learning algorithms. This nonlinear revolution drastically changed the field of statistical learning, and some new research directions such as bioinformatics and data mining were emerged. However, these nonlinear algorithms were mainly based on gradient descent, greedy heuristics and other numerical optimization techniques so suffered from local minima and others. Because their statistical behavior was not well understood, they also experienced overfitting. A third stage in the evolution of statistical learning theory took place in the mid-1990s with the introduction of support vector machine [47] and other kernel-based learning algorithms [49] such as kernel principal component analysis [50], kernel Fisher discriminant analysis [51] and kernel independent component analysis [52]. The new algorithms offered efficiency in analyzing nonlinear relations from computational, statistical and conceptual points of view, and made it possible to do so in the high-dimensional feature space without the dangers of overfitting. The problems of local minima and overfitting that were typical of neural networks and decision trees have been overcome.
The RKHS plays a crucial role in the kernel-based learning algorithms. It follows from the Mercer’s theorem Eq. (1–14) that any symmetric positive definite function \( \kappa(x, y) \) can be rewritten as an inner product between two vectors in the feature space, i.e.,

\[
\kappa(x, y) = \langle \Phi(x), \Phi(y) \rangle
\]

(1–27)

\[
\Phi : x \mapsto \sqrt{\lambda} \varphi_k(x), \quad k = 1, 2, \ldots
\]

There are some different kernels used in statistical learning theory. For example, among others there are

- Gaussian kernel: \( \kappa(x, y) = \frac{1}{\sqrt{2\pi}\sigma} \exp\left\{ -\frac{(x-y)^2}{2\sigma^2} \right\} \), where \( \sigma \) is the kernel width.
- Polynomial kernel: \( \kappa(x, y) = (1 + x \cdot y)^d \), where \( d \) is the polynomial power.
- Sigmoid kernel: \( \tanh(\kappa(x, y) + \beta) \), where \( \beta \) is specified a priori.

Kernel-based learning algorithms use the above idea to map the data from the original input space to a high-dimensional, possibly infinite-dimensional, feature space. By the Moore-Aronszajn Theorem in the previous section, there exists a unique RKHS corresponding to the symmetric positive definite kernel \( \kappa(x, y) \). Therefore the feature space where the transformed data reside is a reproducing kernel Hilbert space, where the nonlinear mapping \( \Phi \) constitutes the basis. Instead of considering the given learning problems in input space, one can deal with the transformed data \( \{\Phi_k(x), k = 1, 2, \ldots\} \) in feature space. When the learning algorithms can be expressed in terms of inner products, this nonlinear mapping becomes particularly interesting and useful since one can employ the kernel trick to compute the inner products in the feature space via kernel functions without knowing the exact nonlinear mapping. The essence of kernel-based learning algorithm is that the inner product of the transformed data can be implicitly computed in the RKHS without explicitly using or even knowing the nonlinear mapping \( \Phi \). Hence, by applying kernels one can elegantly build a nonlinear version of a linear algorithm based on inner products. One of the rationales to non-linearly mapping the data into a high-dimension RKHS is Cover’s theorem on the separability of patterns [53]. Cover’s
theorem, in qualitative terms, states that a complex statistical learning problems cast in a high-dimensional space nonlinearly is more likely to be linearly separable than in a low-dimensional space. By transforming the data into this high-dimensional RKHS and constructing optimal linear algorithms in that space, the kernel-based learning algorithms effectively perform optimal nonlinear pattern recognitions in input space to achieve better separation, estimation, regression and etc.

The research on kernel-based learning algorithms became very active since Vapnik’s seminal paper on support vector machines was published in 1990s. People started to kernelized most the linear algorithms which can be expressed in terms of inner product. One of the drawbacks of the kernel-based learning algorithms is the computational complexity issue. Most kernel-based learning algorithms will eventually result in operations on Gram matrix whose dimension depends on the number of data. For instance, computation of eigenvalues and eigenvectors for a thousand dimensional Gram matrix demands a great deal of computational complexity. Therefore, a great amount of optimization algorithms have been developed to address this issue based on numerical linear algebra. On the other hand, since kernel-based learning from data usually ends up an ill-posed problem, regularization through nonlinear functionals becomes necessary and mandatory. Hence, cross validation is needed to choose an optimal regularization parameter [54].

1.4 A Brief Review of Information-Theoretic Learning (ITL)

In parallel to the developments in kernel-based methods research, independently a research topic called information-theoretic learning (ITL) has emerged [55], where kernel-based density estimators form the essence of this learning paradigm. Information-theoretic learning is a signal processing technique that combines information theory and adaptive systems to implement information filtering without requiring a model of the data distributions. ITL uses the concepts of Parzen windowing applied to Renyi’s entropy definition to obtain a sample by sample algorithm that estimates entropy directly from pairs of sample interactions. By utilizing Renyi’s measure of entropy and approximations
to the Kullback-Leibler probability density divergence, ITL is able to extract information beyond second-order statistics directly from data in a non-parametric manner.

Information-theoretic learning has achieved excellent results on a number of learning scenarios, e.g. blind source separation [56], supervised learning [57] and others [55].

One of the most commonly used cost functions in information-theoretic learning is the quadratic Renyi’s entropy [58]. Renyi’s entropy is a generalization of Shannon’s entropy. Given a PDF $f(x)$ for a random variable $x$, the quadratic Renyi’s entropy is defined as

$$H(x) = -\log \int f^2(x)dx = -\log E[f(x)].$$

Since logarithm function is monotonic, the quantity of interest in adaptive filtering is its argument

$$I(x) = \int f^2(x)dx,$$

which is called information potential, so named due to a similarity with the potential energy field in physics [55]. The concept and properties of information potential have been mathematically studied and a new criterion based on information potential has been proposed, called the MEE (Minimization Error Entropy), to adapt linear and nonlinear systems [59]. MEE serves as an alternative to the conventional MSE (Mean Square Error) in nonlinear filtering with several advantages in performance.

A non-parametric asymptotically unbiased and consistent estimator for a given PDF $f(x)$ is defined as [60]

$$\hat{f}(x) = \frac{1}{N} \sum_{i=1}^{N} \kappa(x, x_i),$$

where $\kappa(\cdot, \cdot)$ is called the Parzen window, or kernel, which is the same symmetric non-negative definite function used in kernel-based learning theory such as Gaussian kernel, polynomial kernel and others [61]. Then by approximating the expectation by the sample mean in Eq. (1–28), we can estimate the information potential directly from the
where \( \{x_i\}_{i=1}^N \) is the data sample and \( N \) is the total number. According to the Mercer’s theorem [13], any symmetric non-negative definite kernel function has an eigen-decomposition as \( \kappa(x,y) = \langle \Phi(x), \Phi(y) \rangle_{\mathcal{H}_\kappa} \), where \( \Phi(x) \) is the nonlinearly transformed data in the RKHS \( \mathcal{H}_\kappa \) induced by the kernel function and the inner product is performed in \( \mathcal{H}_\kappa \). Therefore, we can re-write the estimate of information potential as

\[
\hat{I}(x) = \frac{1}{N^2} \sum_{i=1}^{N} \sum_{j=1}^{N} \kappa(x_i, x_j),
\]

However, the RKHS \( \mathcal{H}_\kappa \) is data independent since the kernel is pre-designed regardless of the data. Therefore, only the estimate of the information potential, not the information potential itself, can be formulated as such. Statistical inference in the RKHS \( \mathcal{H}_\kappa \) does not yield intrinsic geometric interpretation to the statistical information of signals required by ITL.

ITL has also been used to characterize the divergence between two random variables. In information theory, mutual information is one of the quantities that quantifies the divergence between two random variables. Another well-known divergence measure is the Kullback-Leibler divergence [62]. However, the Kullback-Leibler measure is difficult to evaluate in practice without imposing simplifying assumptions about the data, therefore numerical methods are required to evaluate the integrals. In order to integrate the non-parametric PDF estimation via Parzen windowing to provide an efficient estimate, two divergence measures for random variables based on Euclidean difference of vectors inequality and Cauchy-Schwartz inequality respectively have been proposed [55].
The divergence measure based on Euclidean inequality is defined as

\[ D_{ED}(f, g) = \int (f(x) - g(x))^2 \, dx \]
\[ = \int f(x)^2 \, dx - 2 \int f(x)g(x) \, dx + \int g(x)^2 \, dx. \]  

(1–31)

The divergence measure based on Cauchy-Schwartz inequality is given by

\[ D_{CS}(f, g) = -\log \frac{\int f(x)g(x) \, dx}{\sqrt{\left( \int f^2(x) \, dx \right) \left( \int g^2(x) \, dx \right)}}. \]  

(1–32)

Notice that both \( D_{ED}(f, g) \) and \( D_{CS}(f, g) \) are greater than zero, and the equality holds if and only if \( f(x) = g(x) \).

The Euclidean Eq. (1–31) and Cauchy-Schwartz divergence measures Eq. (1–32) can be easily extended to two-dimensional random variables. As a special case, if we substitute the marginal PDFs \( f \) and \( g \) in Eq. (1–31) and Eq. (1–32) by a joint PDF \( f_{1,2}(x_1, x_2) \) and the product of marginal PDFs \( f_1(x_1)f_2(x_2) \) respectively, the Euclidian quadratic mutual information is given by [55]

\[ I_{ED}(f_1, f_2) = -2 \int \int f_{1,2}(x_1, x_2)f_1(x_1)f_2(x_2) \, dx_1 \, dx_2 
+ \int \int f_{1,2}^2(x_1, x_2) \, dx_1 \, dx_2 + \int \int f_1^2(x_1)f_2^2(x_2) \, dx_1 \, dx_2, \]  

(1–33)

and the Cauchy-Schwartz quadratic mutual information is defined as [55]

\[ I_{CS}(f_1, f_2) = -\log \frac{\int \int f_{1,2}(x_1, x_2)f_1(x_1)f_2(x_2) \, dx_1 \, dx_2}{\sqrt{\left( \int \int f_{1,2}^2(x_1, x_2) \, dx_1 \, dx_2 \right) \left( \int \int f_1^2(x_1)f_2^2(x_2) \, dx_1 \, dx_2 \right)}}. \]  

(1–34)

As can be seen from above that \( I_{ED}(f_1, f_2) \geq 0 \) and \( I_{CS}(f_1, f_2) \geq 0 \). If and only if the two random variables are statistically independent, then \( I_{ED}(f_1, f_2) = 0 \) and \( I_{CS}(f_1, f_2) = 0 \). Basically, the Euclidean quadratic mutual information measures the Euclidean difference between the joint PDF and the factorized marginals, and likewise for the Cauchy-Schwartz
quadratic mutual information. Hence minimization of these two measures leads to minimization of mutual information between two random variables. Cauchy-Schwartz divergence measure has been applied to independent component analysis [63], and clustering [64].

One of the limitations of ITL is that it does not convey the time structure of signals because it assumes i.i.d. data. However, in practice most of signals in engineering have correlation in time or temporal structures. It would be helpful to incorporate the temporal structures while still containing high order statistics, for instance working with coded source signals in digital communications.

1.5 Recent Progress on Correntropy

From the previous two brief introductions on RKHS in statistical signal processing and statistical learning algorithms, we notice that there are two different operators, the expectation on random processes and the positive definite kernel on static data, which uniquely determine two different reproducing kernel Hilbert spaces. The expectation operator is data dependent because it operates on the random processes and hence the RKHS induced by the correlation function is embedded with statistics. While the kernel operator in statistical learning algorithms is data independent, which is specified by the designer to be one of the Gaussian kernel, polynomial kernel, sigmoid kernel or others, hence the RKHS induced by one of them in statistical learning only depends on the specific kernel and does not contain the statistical information of data. The conventional statistical signal processing from RKHS perspective induced by the correlation function provides new understanding of second order random processes, however it does not offer new results because all the conventional statistical signal processing had already been carried out without using RKHS tool. On the other hand, the application of data independent kernel in statistical learning theory requires regularization to make the solution unique because the all the learning algorithms evolves the computation of Gram matrix whose dimension is the same as the number of data samples.
One natural question to ask is to combine these two reproducing kernel Hilbert spaces together in some way by means of proposing a new operator such that it is composed of the expectation operator and the pre-designed kernel operator. If successful, we might address the non-linearity in statistical signal processing because the pre-designed kernel can non-linearly map the random processes into a high-dimensional RKHS. Moreover, the pre-designed kernels contain beyond second-order operations on the variables, which might also provide a new tool on non-Gaussian statistical signal processing. The new operator will also capture the time structure of signal because it has the same spirit of conventional autocorrelation, while it might still preserve the higher order statistics information. This can overcome the limitation of Information-Theoretic Learning. The problem of regularization can also be implicitly solved since the new operator employs the expectation.

Recently a new generalized correlation function, called correntropy, has been proposed to combine these two kernels to characterize both the temporal structure and statistical information of random processes [65]. The correntropy has been applied to various signal processing and machine learning problems and produced promising results. The correntropy based matched filter outperforms the conventional matched filter in impulse noise scenario [66]. The correntropy MACE filter has been proposed for image recognition [67]. Since correntropy induces a new RKHS, it is able to bring nonlinearity into the traditional statistical signal processing. Correntropy Wiener filter non-linearly transforms the original random process into the high dimensional RKHS induced by the kernel function while minimizes the mean square error between the desired and output signals. The output signal has been represented by the inner product between those non-linearly transformed input signal and the weights in RKHS. The correntropy Wiener filter exhibits much better performance than the conventional Wiener filter and multilayer perceptron [68]. These up-to-date developments of correntropy clearly demonstrate the promising features and applicable areas, which effectively introduce a new nonlinear signal processing...
paradigm based on reproducing kernel Hilbert space. Unlike the recent advance in kernel-based learning in computer science field, correntropy defines a new data-dependent RKHS and adapts to the intrinsic structure of data.

1.6 Study Objectives

In this dissertation, we analyze the newly proposed correntropy function [65], and present another generalized covariance function, called centered correntropy. The correntropy and centered correntropy functions are typically a combination of expectation operator and pre-designed operator. It can be easily seen that the new operators are also symmetric positive definite and thus induces another reproducing kernel Hilbert spaces which drastically change the structure of the reproducing kernel Hilbert spaces induced by conventional autocorrelation function and the pre-designed kernel function. Although the correntropy and centered correntropy have been applied to some different signal processing and machine learning problems, further theoretical analysis and experimental work are needed to fully elucidate the new concept and evaluate the associated properties. This dissertation strives to serve as one of these efforts.

This dissertation is organized as follows. In chapter 3, the definitions of generalized correlation and covariance functions, which are called correntropy and centered correntropy respectively, are proposed and analyzed from time-domain and frequency-domain. Chapter 4 addresses the geometric structure of the reproducing kernel Hilbert spaces induced by the centered correntropy. A new explicit construction of RKHS with Gaussian kernel is presented. A parametric correntropy function is proposed in chapter 5 to quantify dependence measure. Application of centered correntropy in principal component analysis is presented in Chapter 6. I also apply correntropy in pitch determination in chapter 7 and nonlinear coupling measure in chapter 8. I conclude the work and present some future work in chapter 9.
CHAPTER 2
AN RKHS FRAMEWORK FOR ITL

In this chapter, we propose a reproducing kernel Hilbert space (RKHS) framework for the information-theoretic learning (ITL). The RKHS is uniquely determined by the symmetric non-negative definite kernel function which is defined as the cross information potential (CIP) in ITL. The cross information potential as an integral of product of two probability density functions characterizes similarity between two random variables. We also prove the existence of a one-to-one congruence mapping between the presented RKHS and the Hilbert space spanned by probability density functions. All the cost functions in the original information-theoretic learning formulation can be re-written as algebraic computations on functional vectors in the reproducing kernel Hilbert space. We prove a lower bound for the information potential based on the presented RKHS. The proposed RKHS framework offers an elegant and insightful geometric perspective towards information-theoretic learning.

From the definitions of various cost functions in information-theoretic learning, we see that the most fundamental quantity is the integral of product of two probability density functions (PDFs) $\int f(x)g(x)dx$ which is called the cross information potential (CIP) [55]. Cross information potential measures the similarity between two PDFs, while the information potential Eq. (1–28) is nothing but a measure of self-similarity. CIP appears both in Euclidean and Cauchy-Schwartz divergence measures. In this chapter, we shall develop the reproducing kernel Hilbert space framework for information-theoretic learning based on the cross information potential.

2.1 The RKHS based on ITL

The RKHS framework based on the PDFs of the data for ITL is proposed in this section. We first focus on the development for one-dimensional case. The extension to multi-dimension is straightforward. We form a $L_2$ space consisting of all one-dimensional PDFs, and define an inner product in $L_2$. Since the inner product is symmetric non-negative
definite, it uniquely determines a reproducing kernel Hilbert space $\mathcal{H}_V$. We then prove that the inner product itself is indeed a reproducing kernel in $\mathcal{H}_V$.

### 2.1.1 The $L_2$ Space of PDFs

Let $\mathcal{E}$ be the set that consists of all square integrable one-dimensional probability density functions, i.e., $f_i(x) \in \mathcal{E}$, $\forall i \in \mathbb{I}$, where $\int f_i(x)^2 \, dx < \infty$ and $\mathbb{I}$ is an index set. We then form a linear manifold

$$\left\{ \sum_{i \in I} \alpha_i f_i(x) \right\}$$

(2–1)

for any $I \subset \mathbb{I}$ and $\alpha_i \in \mathbb{R}$. Close the set in Eq. (2–1) topologically according to the convergence in the mean using the norm

$$\|f_i(x) - f_j(x)\| = \sqrt{\int (f_i(x) - f_j(x))^2 \, dx}, \quad \forall i, j \in \mathbb{I}$$

(2–2)

and denote the set of all linear combinations of PDFs and its limit points by $L_2(\mathcal{E})$. $L_2(\mathcal{E})$ is an $L_2$ space on PDFs. Moreover, by the theory of quadratically integrable functions, we know that the linear space $L_2(\mathcal{E})$ forms a Hilbert space if an inner product is imposed accordingly. Given any two PDFs $f_i(x)$ and $f_j(x)$ in $\mathcal{E}$, we can define an inner product as

$$\langle f_i(x), f_j(x) \rangle_{L_2} = \int f_i(x)f_j(x) \, dx, \quad \forall i, j \in \mathbb{I}. \quad (2–3)$$

Notice that this inner product is exactly the cross information potential $[55]$. This definition of inner product has a corresponding norm of Eq. (2–2). Hence, the $L_2(\mathcal{E})$ equipped with the inner product Eq. (2–3) is a Hilbert space. However, it is not a reproducing kernel Hilbert space because the inner product does not satisfy the reproducing property in $L_2(\mathcal{E})$. Next we show that the inner product Eq. (2–3) is symmetric non-negative definite, and by the Moore-Aronszajn theorem it uniquely determines a reproducing kernel Hilbert space.
2.1.2 RKHS $H_V$ Based on $L_2(\mathcal{E})$

First, we define a bivariate function on the set $\mathcal{E}$ as

$$V(f_i, f_j) = \int f_i(x)f_j(x)dx, \quad \forall i, j \in I.$$ \hspace{1cm} (2–4)

This function is also the definition of the inner product Eq. (2–3), and the cross information potential between two PDFs. This will be the kernel function in the RKHS $H_V$ constructed below. In reproducing kernel Hilbert space theory, the kernel function is a measure of similarity between functionals. As pointed out earlier, the cross information potential is a similarity measure between two probability density functions, hence it is natural and meaningful to define the kernel function as such. Next, we show that function Eq. (2–4) is symmetric non-negative definite in $\mathcal{E}$.

**Property 1 (Non-Negative Definiteness):** The function Eq. (2–4) is symmetric non-negative definite in $\mathcal{E} \times \mathcal{E} \rightarrow \mathcal{R}$.

**Proof:** The symmetry is obvious. Given any positive integer $N$, any set of \{f_1(x), f_2(x), \ldots f_N(x)\} $\in \mathcal{E}$ and any not all zero real numbers $\{\alpha_1, \alpha_2, \ldots, \alpha_N\}$, by definition we have

$$\sum_{i=1}^{N} \sum_{j=1}^{N} \alpha_i \alpha_j V(f_i, f_j) = \sum_{i=1}^{N} \sum_{j=1}^{N} \alpha_i \alpha_j \int f_i(x) f_j(x) dx$$

$$= \int \left( \sum_{i=1}^{N} \alpha_i f_i(x) \right) \left( \sum_{j=1}^{N} \alpha_j f_j(x) \right) dx = \int \left( \sum_{i=1}^{N} \alpha_i f_i(x) \right)^2 dx \geq 0.$$ 

Hence, $V(f_i, f_j)$ is symmetric non-negative definite, and it is also a kernel function. ■

According to the Moore-Aronszajn theorem, there is a unique reproducing kernel Hilbert space, denoted by $H_V$, associated with the symmetric non-negative definite function Eq. (2–4). We construct the RKHS $H_V$ from bottom-up. Since function Eq. (2–4) is symmetric and non-negative definite, it also has an eigen-decomposition by the Mercer’s theorem [13] as

$$V(f_i, f_j) = \sum_{k=1}^{\infty} \lambda_k \psi_k(f_i) \psi_k(f_j),$$ \hspace{1cm} (2–5)
where \( \{ \psi_k(f), k = 1, 2, \ldots \} \) and \( \{ \lambda_k, k = 1, 2, \ldots \} \) are sequences of eigenfunctions and corresponding eigenvalues of the kernel function \( V(f, f_j) \) respectively. The series above converges absolutely and uniformly on \( \mathcal{E} \times \mathcal{E} \) [13].

Then we define a space \( \mathcal{H}_V \) consisting of all functionals \( \mathcal{G}(\cdot) \) whose evaluation for any given PDF \( f_i(x) \in \mathcal{E} \) is defined as

\[
\mathcal{G}(f_i) = \sum_{k=1}^{\infty} \lambda_k a_k \psi_k(f_i), \tag{2-6}
\]

where the sequence \( \{a_k, k = 1, 2, \ldots \} \) satisfies the following condition

\[
\sum_{k=0}^{\infty} \lambda_k a_k^2 < \infty. \tag{2-7}
\]

Furthermore we define an inner product of two functionals in \( \mathcal{H}_V \) as

\[
\langle \mathcal{G}, \mathcal{F} \rangle_{\mathcal{H}_V} = \sum_{k=0}^{\infty} \lambda_k a_k b_k, \tag{2-8}
\]

where \( \mathcal{G} \) and \( \mathcal{F} \) are of form Eq. (2–6), and \( a_k \) and \( b_k \) satisfy property Eq. (2–7).

It can be verified that the space \( \mathcal{H}_V \) equipped with the kernel function Eq. (2–4) is indeed a reproducing kernel Hilbert space and the kernel function \( V(f_i, \cdot) \) is a reproducing kernel because of the following two properties:

1. \( V(f_i, f_j) \) as a function of \( f_i(x) \) belongs to \( \mathcal{H}_V \) for any given \( f_j(x) \in \mathcal{E} \) because we can rewrite \( V(f_i, f_j) \) as

\[
V(f_i, \cdot)(f_j) = \sum_{k=1}^{\infty} \lambda_k b_k \psi_k(f_j), \quad b_k = \psi_k(f_i).
\]

That is, the constants \( \{b_k, k = 1, 2, \ldots \} \) become the eigenfunctions \( \{\psi_k(f_i), k = 1, 2, \ldots \} \) in the definition of \( \mathcal{G} \). Therefore,

\[
V(f_i, \cdot) \in \mathcal{H}_V, \quad \forall f_i(x) \in \mathcal{E}.
\]
2. Given any $G \in \mathcal{H}_V$, the inner product between the reproducing kernel and $G$ yields the function itself by the definition Eq. (2–8)

$$\langle G, V(f_i, \cdot) \rangle_{\mathcal{H}_V} = \sum_{k=0}^{\infty} \lambda_k a_k b_k = \sum_{k=1}^{\infty} \lambda_k a_k \psi_k(f_i) = G(f_i).$$

This is so called the reproducing property.

Therefore, $\mathcal{H}_V$ is a reproducing kernel Hilbert space with the kernel function and inner product defined above. ■

By the reproducing property, we can re-write the kernel function Eq. (2–5) as

$$V(f_i, f_j) = \langle V(f_i, \cdot), V(f_j, \cdot) \rangle_{\mathcal{H}_V} \quad (2-9)$$

$$V(f_i, \cdot) : f_i \mapsto \sqrt{\lambda_k \psi_k(f_i)}, \quad k = 1, 2, \ldots$$

The reproducing kernel nonlinearly maps the original PDF $f_i(x)$ into the RKHS $\mathcal{H}_V$.

We emphasize here that the reproducing kernel $V(f_i, f_j)$ is data-dependent by which we mean the norm of nonlinearly transformed vector in the RKHS $\mathcal{H}_V$ is dependent on the PDF of the original random variable because

$$\|V(f_i, \cdot)\|^2 = \langle V(f_i, \cdot), V(f_i, \cdot) \rangle_{\mathcal{H}_V} = \int f_i(x)^2 dx.$$ 

This is very different from the reproducing kernel $\kappa(x, y)$ used in kernel-based learning theory. The norm of nonlinearly projected vector in the RKHS $\mathcal{H}_\kappa$ does not rely on the statistical information of the original data since

$$\|\Phi(x)\|^2 = \langle \Phi(x), \Phi(x) \rangle_{\mathcal{H}_\kappa} = \kappa(0)$$

if we use translation-invariant kernel function [61]. The value of $\kappa(0)$ is a constant regardless of the original data. Consequently, the reproducing kernel Hilbert spaces $\mathcal{H}_V$ and $\mathcal{H}_\kappa$ determined by $V(f_i, f_j)$ and $\kappa(x, y)$ respectively are very different in nature.
2.1.3 Congruence Map Between $\mathcal{H}_V$ and $L_2(\mathcal{E})$

We have presented two Hilbert spaces, the Hilbert space $L_2(\mathcal{E})$ of PDFs and the reproducing kernel Hilbert space $\mathcal{H}_V$. Even though their elements are very different, there actually exists a one-to-one congruence mapping $\Psi$ (isometric isomorphism) from RKHS $\mathcal{H}_V$ onto $L_2(\mathcal{E})$ such that

$$\Psi(V(f_i, \cdot)) = f_i(x).$$

(2–10)

Notice that the mapping $\Psi$ preserves isometry between $\mathcal{H}_V$ and $L_2(\mathcal{E})$ since by definitions of inner product Eq. (2–3) in $L_2(\mathcal{E})$ and Eq. (2–9) in $L_2(\mathcal{E})$

$$\langle V(f_i, \cdot), V(f_j, \cdot) \rangle_{\mathcal{H}_V} = \langle f_i(x), f_j(x) \rangle_{L_2} = \langle \Psi(V(f_i, \cdot)), \Psi(V(f_j, \cdot)) \rangle_{L_2}.$$  

That is, the mapping $\Psi$ maintains the inner products in both $\mathcal{H}_V$ and $L_2(\mathcal{E})$.

In order to obtain an explicit representation of $\Psi$, we define an orthogonal function sequence $\{\xi_m(x), m = 1, 2, \ldots\}$ satisfying

$$\int \xi_k(x)\xi_m(x)dx = \begin{cases} 0, & k \neq m \\ \lambda_k, & k = m \end{cases}$$

and

$$\int \sum_{k=1}^{\infty} \psi_k(f_i)\xi_k(x)dx = 1,$$  

(2–11)

where $\lambda_k$ and $\psi_k(f_i)$ are eigenvalue and eigenfunction associated with the kernel function $V(f_i, f_j)$ by the Mercer’s theorem Eq. (2–5). We achieve an orthogonal decomposition of the probability density function as

$$f(x) = \sum_{k=1}^{\infty} \psi_k(f)\xi_k(x), \quad \forall f(x) \in \mathcal{E}.$$  

(2–12)

The normality condition is fulfilled by the assumption Eq. (2–11).
Note that the congruence map $\Psi$ can be characterized as the unique mapping from $\mathcal{H}_V$ into $L_2(\mathcal{E})$ satisfying the condition that for every functional $\mathcal{G}$ in $\mathcal{H}_V$ and every $j$ in $\mathbb{I}$

$$\int \Psi(\mathcal{G}) f_j(x) dx = \langle \mathcal{G}, V(f_j, \cdot) \rangle_{\mathcal{H}_V} = \mathcal{G}(f_j).$$

(2–13)

It is obvious that $\Psi$ in Eq. (2–10) fulfills the condition Eq. (2–13). Then the congruence map can be represented explicitly as

$$\Psi(\mathcal{G}) = \sum_{k=1}^{\infty} a_k \xi_k(x), \quad \forall \mathcal{G} \in \mathcal{H}_V,$$

(2–14)

where $a_k$ satisfies condition Eq. (2–7).

To prove the representation Eq. (2–14) is a valid and unique map, substituting Eq. (2–12) and Eq. (2–14) into Eq. (2–13), we obtain

$$\int \sum_{k=1}^{\infty} a_k \xi_k(x) \sum_{m=1}^{\infty} \psi_m(f_j) \xi_m(x) dx = \sum_{k=1}^{\infty} \sum_{m=1}^{\infty} a_k \psi_m(f_j) \int \xi_k(x) \xi_m(x) dx = \sum_{k=1}^{\infty} \lambda_k a_k \psi_k(f_j) = \mathcal{G}(f_j).$$

In summary, we provide an explicit representation for the congruence map $\Psi$ from RKHS $\mathcal{H}_V$ into $L_2(\mathcal{E})$. These two spaces are equivalent in some geometrical sense. However it should be emphasized that the constituting elements are very different in nature. The RKHS isometry framework offers a natural link between stochastic and deterministic functional analysis. Hence, it is more appealing to use RKHS $\mathcal{H}_V$ for information-theoretic learning as we will show in next section.

2.1.4 Extension to Multi-dimensional PDFs

Extension of $\mathcal{H}_V$ to multi-dimensional PDFs is straightforward since the definitions and derivations in the previous section can be easily adapted into multi-dimensional probability density functions. Now let $\mathcal{E}_m$ be the set that consists of all square integrable $m$ dimensional probability density functions, i.e., $f_{i,m}(x_1, \ldots, x_m) \in \mathcal{E}_m, \forall i \in \mathbb{I}$ and $m \in \mathbb{N}$,
where $\int f_{i,m}(x_1, \ldots, x_m)^2dx_1, \ldots, dx_m < \infty$ and $\mathbb{I}$ is the index set. We need to change the definition of kernel function Eq. (2–4) to
\[
V(f_{i,m}, f_{j,m}) = \int f_{i,m}(x_1, \ldots, x_m)f_{j,m}(x_1, \ldots, x_m)dx_1, \ldots, dx_m, \quad \forall i, j \in \mathbb{I}.
\]
Then every definitions and derivations might as well be modified accordingly in the previous section. Let $\mathcal{H}_V(m)$ denote the reproducing kernel Hilbert space determined by the kernel function for $m$ dimensional PDFs. The proposed RKHS framework is consistent with dimensionality of PDFs.

### 2.2 ITL Cost Functions in RKHS Framework

In this section, we re-examine the ITL cost functions in the proposed RKHS framework.

First, as the kernel function $V(f_i, f_j)$ in $\mathcal{H}_V$ is defined as the cross information potential between two PDFs, immediately we have
\[
\int f(x)g(x)dx = \langle V(f, \cdot), V(g, \cdot) \rangle_{\mathcal{H}_V}. \tag{2–15}
\]
That is, the cross information potential is the inner product between two nonlinearly transformed functionals in the RKHS $\mathcal{H}_V$. The inner product quantifies similarity between two functionals which is consistent with the definition of cross information potential. The information potential can thus be specified as the inner product of a functional with respect to itself
\[
\int f(x)^2dx = \langle V(f, \cdot), V(f, \cdot) \rangle_{\mathcal{H}_V} = \|V(f, \cdot)\|^2. \tag{2–16}
\]
The information potential appears as the norm square of nonlinearly transformed functional in the RKHS $\mathcal{H}_V$. Therefore, minimizing error entropy in ITL turns out to be maximization of norm square in the RKHS $\mathcal{H}_V$ (because the information potential is the argument of the $– \log$ in Renyi’s quadratic entropy). As stated in ITL, MEE employs higher-order statistics in nonlinear adaptive systems training since it is based on Renyi’s quadratic entropy [59]. We observe here that the higher-order statistics in MEE becomes
second norm in the RKHS $H_V$. However, the nonlinearly transformed functional $V(f, \cdot)$ is deterministic in $H_V$. Hence, the proposed RKHS framework provides a link between stochastic and deterministic transformation. The conventional mean square error has also been re-written as norm square of projected vectors in the RKHS $H_R$ induced by the covariance function [25]. But the RKHS $H_R$ only takes the second-order statistics into account, i.e., the mean square error. The RKHS $H_V$ implicitly embeds higher-order statistics. Compared to the RKHS $H_\kappa$ induced by the pre-designed kernel function used in the machine learning, our framework is more elegant theoretically because it corresponds to the definition of information potential directly without employing any kernel-based PDF estimator. From the computational point of view, the estimate of information potential based on Parzen window PDF estimator yields a direct calculation of information quantity from data. However from the theoretical perspective, it is more appropriate to define the RKHS framework based on the information potential itself instead of the estimate using a kernel-based PDF estimator.

Based on the reformulations of cross information potential Eq. (2–15) and information potential Eq. (2–16) in RKHS $H_V$, we are ready to re-write the one-dimensional Euclidean Eq. (1–31) and Cauchy-Schwartz divergence measures Eq. (1–32) in terms of operations on functionals in $H_V$. First,

$$D_{ED}(f, g) = \|V(f, \cdot) - V(g, \cdot)\|^2.$$  

That is, the Euclidean divergence measure is in fact the norm square of difference between two corresponding functionals in $H_V$. This interpretation resembles the conventional definition of Euclidean distance more than the original description Eq. (1–31) does. The Cauchy-Schwartz divergence measure can be phrased as

$$D_{CS}(f, g) = -\log \frac{\langle V(f, \cdot), V(g, \cdot) \rangle_{H_V}}{\|V(f, \cdot)\| \cdot \|V(g, \cdot)\|} = -\log(\cos \theta),$$
where $\theta$ is the angle between two functional vectors $V(f, \cdot)$ and $V(g, \cdot)$. Therefore, the Cauchy-Schwartz divergence measure truly depicts the separation of two functional vectors in the RKHS $\mathcal{H}_V$. When two vectors lie in the same direction and the angle $\theta = 0^\circ$, $D_{CS}(f, g) = 0$. If two vectors are perpendicular to each other ($\theta = 90^\circ$), $D_{CS}(f, g) = \infty$. The RKHS $\mathcal{H}_V$ supplies rich geometric insights into the original definitions of the two divergence measures.

To extend the same formulation to the Euclidean and Cauchy-Schwartz quadratic mutual information Eq. (1–33) and Eq. (1–34), consider the product of marginal PDFs $f_1(x_1)f_2(x_2)$ as a special subset $\mathcal{A}_2$ of the 2-dimensional square integrable PDFs set $\mathcal{E}_2$ where the joint PDF can be factorized into product of marginals, i.e., $\mathcal{A}_2 \subseteq \mathcal{E}_2$. Then both measures characterize different geometric information between the joint PDF and the factorized marginal PDFs. The Euclidean quadratic mutual information Eq. (1–33) can be expressed as

$$I_{ED}(f_1, f_2) = \|V(f_{1,2}, \cdot) - V(f_1f_2, \cdot)\|^2,$$

where $V(f_{1,2}, \cdot)$ is the functional in $\mathcal{H}_{V(2)}$ corresponding to the joint PDF $f_{1,2}(x_1, x_2)$, and $V(f_1f_2, \cdot)$ is for the product of the marginal PDFs $f_1(x_1)f_2(x_2)$. Similarly, the Cauchy-Schwartz quadratic mutual information can be re-written as

$$I_{CS}(f_1, f_2) = -\log \langle V(f_{1,2}, \cdot), V(f_1f_2, \cdot) \rangle_{\mathcal{H}_V} = -\log(\cos \gamma). \quad (2–17)$$

The angle $\gamma$ is the separation between two functional vectors in $\mathcal{H}_{V(2)}$. When two random variables are independent ($f_{1,2}(x_1, x_2) = f_1(x_1)f_2(x_2)$ and $\mathcal{A}_2 = \mathcal{E}_2$), $\gamma = 0^\circ$ and the divergence measure $I_{CS}(f_1, f_2) = 0$ since two sets are equal. If $\gamma = 90^\circ$, two vectors in $\mathcal{H}_{V(2)}$ are orthogonal and the joint PDF is singular to the product of marginals. In this case, the divergence measure is infinity.

The proposed RKHS framework provides an elegant and insightful geometric perspective towards information-theoretic learning. All the cost functions in ITL can now be re-expressed in terms of algebraic operations on functionals in RKHS $\mathcal{H}_V$.  

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2.3 A Lower Bound for the Information Potential

Based on the proposed RKHS framework for the information-theoretic learning, we derive a lower bound for the information potential Eq. (1–28) in this section. First we cite the projection theorem in Hilbert space that we will use in the following proof.

**Theorem 2 (Projection in Hilbert Space)** Let \( H \) be a Hilbert space, \( M \) be a Hilbert subspace of \( H \) spanned by \( N \) linearly independent vectors \( u_1, u_2, \ldots, u_N \), \( s \) be a vector in \( H \), and \( d \) be a quantity such that

\[
    d = \inf \| s - u \|, \quad \forall \; u \in M.
\]

Then there exists a unique vector, denoted as \( P(s|M) \), in \( M \) such that

\[
P(s|M) = \sum_{i=1}^{N} \sum_{j=1}^{N} \langle s, u_i \rangle K^{-1}(i, j) u_j,
\]

where \( K(i, j) \) is the \( N \times N \) Gram matrix whose \((i, j)\) is given by \( \langle u_i, u_j \rangle \). The projected vector \( P(s|M) \) also satisfies the following conditions:

\[
    \| s - P(s|M) \| = d = \min \| s - u_i \|, \quad (2-19)
\]

\[
    \langle s - P(s|M), u_i \rangle = 0, \quad \forall \; u_i \in M,
\]

\[
    \langle P(s|M), u_i \rangle = \langle s, u_i \rangle, \quad \forall \; u_i \in M. \quad (2-20)
\]

The geometrical explanation of the theorem is straightforward. Readers can refer to [21] for a thorough proof. Now we state the proposition on a lower bound for the information potential.

**Proposition (Lower Bound for the Information Potential)** Let \( V(f, \cdot) \) be a vector in the RKHS \( H_V \) induced by the kernel \( V \), \( M \) be a subspace of \( H_V \) spanned by \( N \) linearly independent vectors \( V(g_1, \cdot), V(g_1, \cdot), \ldots, V(g_N, \cdot) \in H_V \), Then,

\[
    \int f(x)^2 dx = \| V(f, \cdot) \|^2 \geq \sum_{i,j=1}^{N} \langle V(f, \cdot), V(g_i, \cdot) \rangle_{H_V} G^{-1}(i, j) \langle V(f, \cdot), V(g_j, \cdot) \rangle_{H_V},
\]

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where \( G(i,j) \) is the \( N \times N \) Gram matrix whose \( (i,j) \) term is defined as \( \langle V(g_i, \cdot), V(g_j, \cdot) \rangle_{\mathcal{H}_V} \).

**Proof:** first by the projection theorem Eq. (2–18), we can find the orthogonal projection of \( V(f, \cdot) \) onto the subspace \( \mathcal{M} \) as

\[
P(V(f, \cdot)|\mathcal{M}) = \sum_{i,j=1}^{N} \langle V(f, \cdot), V(g_i, \cdot) \rangle_{\mathcal{H}_V} G^{-1}(i,j) V(g_j, \cdot).
\]

Since the Gram matrix is symmetric and positive definite, the inverse always exists. Next, we calculate the norm square of the projected vector by Eq. (2–20),

\[
\|P(V(f, \cdot)|\mathcal{M})\|^2 = \langle V(f, \cdot), P(V(f, \cdot)|\mathcal{M}) \rangle_{\mathcal{H}_V} = \sum_{i,j=1}^{N} \langle V(f, \cdot), V(g_i, \cdot) \rangle_{\mathcal{H}_V} G^{-1}(i,j) \langle V(f, \cdot), V(g_j, \cdot) \rangle_{\mathcal{H}_V}.
\]

On the other hand, the projection residual defined in Eq. (2–19) satisfies

\[
d^2 = \|V(f, \cdot)\|^2 - \|P(V(f, \cdot)|\mathcal{M})\|^2 \geq 0. \tag{2–21}
\]

Combining Eq. (2–21) and Eq. (2–21), we come to the conclusion of our proposition Eq. (2–21). ■

The proposition generalizes the Carmer-Rao inequality in the statistical estimation theory. It can also be viewed as an approximation to the functional norm by a set of orthogonal bases. Equation Eq. (2–21) offers a theoretical lower bound for minimization of information potential.

### 2.4 Discussions

In this section, we relate our work to the concepts of information geometry and probability product kernels.

#### 2.4.1 Non-parametric vs. Parametric

The RKHS framework presented in this article elucidates the geometric structure on the space of all probability density distributions. Since it does not assume any models for the PDFs, it is non-parametric and infinite-dimensional. In statistics, information geometry studies the intrinsic geometry in a finite-dimensional, parametric statistical
manifold formed by all PDFs [69]. Extension to infinite-dimensional, non-parametric sub-manifold has been advanced [70]. For finite-dimensional, parametric families of PDFs, the only invariant metric to the tangent space is the Riemannian structure defined by the Fisher information [69, 71]

\[ g_{ij}(\theta) = \mathbf{E} \left[ \frac{\partial \log f(x; \theta)}{\partial \theta_i} \frac{\partial \log f(x; \theta)}{\partial \theta_j} \right] \]

in the component form. The Riemannian metric coincides infinitesimally with the double of the Kullback-Leibler divergence. More interestingly, the Fisher information is a symmetric non-negative definite function defined in the parameter space. Therefore, it uniquely determines a reproducing kernel Hilbert space. But, it is very different from our approach because the proposed RKHS framework assumes a non-parametric probability density functions. The non-negative definite kernel function Eq. (2–4) is defined directly in the PDF space, however the kernel function in information geometry is defined in the parameter space since it aims at estimating model parameters from data. Hence, both methodologies define non-parametric and parametric reproducing kernel Hilbert spaces respectively to tackle problems of interest from different perspectives.

2.4.2 Kernel Function as a Dependence Measure

The kernel function we defined characterizes relationships between probability density functions. For instance, the one-dimensional kernel function \( V(f, g) \) quantifies how similar one PDF is to the other. The two-dimensional function \( V(f_{1,2}, f_1 f_2) \) specifies the relationship between joint PDF and product of marginal PDFs. Therefore it also measures how dependent one random variable is to the other. The Cauchy-Schwartz quadratic mutual information Eq. (2–17) was applied to independent component analysis based on this interpretation [63]. Using probability distributions to measure similarity is nothing new. One customary quantity is the Kullback-Leibler divergence. However, it is not positive definite nor symmetric, and hence does not have a reproducing kernel Hilbert
space associated with it. Therefore, KL divergence lacks the geometric advantage of RKHS that our kernel function possesses.

Recently, several probability product kernels have been proposed in machine learning field to use ensembles instead of individual data to capture dependence between generative models [72]. It is shown that the Bhattacharyya coefficient defined by

$$\kappa(f_i, f_j) = \int \sqrt{f_i(x)f_j(x)}dx$$

is a reproducing kernel [73]. The expected likelihood kernel in [73] is exactly the cross information potential. But as they proposed the probability product kernel purely from machine learning point of view, it failed to relate to a broader information theoretic framework. Our contribution in this chapter is to independently propose a reproducing kernel Hilbert space framework for information-theoretic learning, construct the RKHS from bottom-up, and prove the validity mathematically. Therefore the kernel function has a rich information-theoretic interpretation. Moreover, as ITL is mainly applied to adaptive signal processing, we employ non-parametric method to compute the cross information potential kernel and other quantities without an explicit probability density function estimation. However, a parametric generative model is assumed in order to calculate the kernel in their approach [72, 73].

2.5 Conclusion

In this chapter, we present a geometric structure for the information-theoretic learning methodology. The proposed reproducing kernel Hilbert space framework is determined by the symmetric non-negative definite kernel function which is defined as the cross information potential. The kernel function quantifies the similarity between two transformed functionals in the RKHS $\mathcal{H}_V$. We can re-write all the cost functions in ITL in terms of algebraic operations on the functionals in the proposed reproducing kernel Hilbert space. These formulations offer a rich geometric for the original information-theoretic learning algorithms. Compared to a previous RKHS framework by the pre-designed
kernel function, ours is built directly on the probability density functions and contains the statistical information of the data. Hence, the RKHS $\mathcal{H}_V$ provides an elegant geometric structure intrinsic to the data statistics. The future work would include deriving least projection theorem in RKHS $\mathcal{H}_V$ so that we might present minimum information potential estimator and others directly from the functionals.
CHAPTER 3
CORRENTROPY AND CENTERED CORRENTROPY FUNCTIONS

3.1 Autocorrentropy and Crosscorrentropy Functions

A random process is completely specified by its $n$-dimensional joint probability distribution function for every positive integer $n$. But in reality, it is not practical, or even possible, to determine all of its $n$-dimensional joint probability distribution functions. Therefore, we might only use the 2-dimensional distribution functions to partially characterize a given random process. In many engineering applications, the 2-dimensional distribution functions offer a substantial information to tackle most the problems. The most important descriptions for the 2-dimensional distributions are the mean, autocorrelation, and autocovariance functions. This way of partial (second-order) characterization provides practical measurements of random processes and also well suits linear operations on random processes [74].

But when the application is required to deal with nonlinear systems or non-gaussian signals, the second order statistics are not sufficient because they fail to capture the nonlinearity and higher order statistics intrinsic to the problems. There have been many attempts to tackle nonlinear and non-Gaussian signal processing. For instance, the Wiener and Hammerstein models are the early methods proposed to implement optimal nonlinear system identification [75]. In these models, a static nonlinearity chosen \textit{a priori} is placed in front of (or after) a linear time-invariant system, and the optimal solution is obtained via Wiener-Hopf equation [75]. Others include Volterra series [76]. Recently Principe \textit{et al} applied information theory quantities, like entropy and mutual information, to adaptive signal processing, which is named information-theoretic learning (ITL) [55]. But ITL methods lack the time structure information of the random processes. To incorporate both the time structure and the high order statistics, a new generalized correlation function, called correntropy, was proposed by Santamaria \textit{et al} recently in [65]. This generalized correlation function can also be viewed as a correlation function for the transformed
random process in a high dimensional reproducing kernel Hilbert space via nonlinear mapping imposed by the kernel. The linear operations on the transformed random process in the feature space will become nonlinear operations on the original random process in the input space by means of inverse nonlinear mapping. Therefore, this generalized correlation function also conveys the nonlinearity in the random processes.

**Definitions:** Given a random process \( \{x_t : t \in \mathbb{T}\} \) with \( t \) typically denoting time and \( \mathbb{T} \) being an index set of interest, the generalized correlation function, named autocorrentropy, is defined as

\[
V(t, s) = E[\kappa(x_t, x_s)],
\]

the generalized covariance function, named centered autocorrentropy, is defined as

\[
U(t, s) = E_{x_t, x_s}[\kappa(x_t, x_s)] - E_{x_t} E_{x_s}[\kappa(x_t, x_s)],
\]

for each \( t \) and \( s \) in \( \mathbb{T} \), where \( E \) denotes the statistical expectation operator and \( \kappa(\cdot, \cdot) \) is a symmetric positive definite kernel function. Notice that the correntropy is the joint expectation of \( \kappa(x_t, x_s) \), while the centered correntropy is the difference between joint expectation and marginal expectation of \( \kappa(x_t, x_s) \).

In the literature, there have been several symmetric positive definite kernel functions proposed in machine learning, function approximation, density estimation, support vector machines, and etc. For example, the sigmoidal kernel, Gaussian kernel, polynomial kernel and spline kernel just to name a few. The mostly used kernel function is the Gaussian kernel which is given by

\[
\kappa(x_t, x_s) = \frac{1}{\sqrt{2\pi\sigma}} \exp \left\{ -\frac{(x_t - x_s)^2}{2\sigma^2} \right\}.
\]

By the Mercer’s theorem in chapter 1 Eq. (1–14), any symmetric positive definite function \( \kappa(x_t, x_s) \) can be rewritten as an inner product between two vectors in the feature
The nonlinear transformation $\Phi$ maps the random process \( \{ x_t : t \in T \} \) into another random process \( \{ \Phi(x_t) : t \in T \} \) in the high-dimensional feature space, which is a reproducing kernel Hilbert space induced by the kernel according to the Moore-Aronzajn theorem Eq. (1–2). Therefore we can rewrite the autocorrentropy in terms of the nonlinear mapping $\Phi$ as

\[
V(t, s) = E[\langle \Phi(x_t), \Phi(x_s) \rangle]. \tag{3–5}
\]

Likewise the centered autocorrentropy can also be expressed as

\[
U(t, s) = E[\langle \Phi(x_t), \Phi(x_s) \rangle] - \langle E[\Phi(x_t)], E[\Phi(x_s)] \rangle = E[\langle \Phi(x_t) - E[\Phi(x_t)], \Phi(x_s) - E[\Phi(x_s)] \rangle]. \tag{3–6}
\]

It can be seen that the correntropy function is a “conventional” correlation function for the transformed random process in the high-dimensional RKHS. While the centered correntropy is nothing but the correntropy for the zero mean (centered) random process \( \{ \Phi(x_t) - E[\Phi(x_t)] : t \in T \} \). This way of defining the generalized correlation and covariance functions is in the same spirit of the standard correlation and covariance functions.

Correntropy can be applied both to one time series, called the autocorrentropy as we have defined above, or a pair of multidimensional random variables, called the crosscorrentropy. The definitions of crosscorrentropy and centered crosscorrentropy functions are straightforward. They compute the generalized correlation across the space structure instead of time structure.

**Definition:** Given two random variables $x$ and $y$, the crosscorrentropy function is defined as

\[
V(x, y) = E[\kappa(x, y)] = E[\langle \Phi(x), \Phi(y) \rangle]. \tag{3–7}
\]
and the centered crosscorrentropy is defined as

\[ U(x, y) = E_{x,y}[\kappa(x, y)] - E_xE_y[\kappa(x, y)] \]
\[ = E[\langle \Phi(x) - E[\Phi(x)], \Phi(y) - E[\Phi(y)] \rangle] \]
\[ = \int \int_{\Omega} \kappa(x, y)(f_{X,Y}(x, y) - f_X(x)f_Y(y))dxdy, \] (3–8)

where \( E \) denotes the statistical expectation operator, \( \kappa(\cdot, \cdot) \) is a symmetric positive definite kernel function, and \( \Phi \) is the nonlinear transformation associated with kernel \( \kappa(\cdot, \cdot) \) Eq. (1–27). Notice that correntropy is the joint expectation of \( \kappa(x, y) \), while the centered correntropy is the difference between the joint expectation and marginal expectation of \( \kappa(x, y) \).

The correntropy and centered correntropy functions for random variables share the same properties as the ones for random processes since they can also be viewed as one instant of correntropy and centered correntropy functions for random processes. Without ambiguity, we will call both autocorrentropy and crosscorrentropy correntropy functions in the followings.

If the random process \( \{x_t : t \in T\} \) is pair-wise independent, in other word,

\[ f_{X_t, X_s}(x_t, x_s) = f_{X_t}(x_t)f_{X_s}(x_s), \quad \forall t \neq s \in T, \] (3–9)

where \( f_{X_t, X_s}(x_t, x_s) \) is the joint probability density function (PDF) and \( f_{X_t}(x_t), f_{X_s}(x_s) \) are the marginal PDFs, then the correntropy at \( t \neq s \) becomes

\[ V(t, s) = \int \int \langle \Phi(x_t), \Phi(x_s) \rangle f_{X_t, X_s}(x_t, x_s)dxdy \]
\[ = \int \int \langle \Phi(x_t), \Phi(x_s) \rangle f_{X_t}(x_t)f_{X_s}(x_s)dxdy \]
\[ = \langle \int \Phi(x_t)f_{X_t}(x_t)dxt, \int \Phi(x_s)f_{X_s}(x_s)ds \rangle \]
\[ = \langle E[\Phi(x_t)], E[\Phi(x_s)] \rangle \]
\[ = E_{x_t}E_{x_s}[\kappa(x_t, x_s)] \approx \frac{1}{N^2} \sum_{i=1}^{N} \sum_{j=1}^{N} \kappa(x_i, x_j) \] (3–10)
We have used the sample mean to estimate the statistical expectation and the independence property in the computation above, where \( \{x_1, x_2, \ldots, x_N\} \) is one realization of the random process. The quantity Eq. (3–10) is called *information potential* and corresponds to the argument of the logarithm of the quadratic Renyi’s entropy when a Parzen window estimator is used [55]. Hence the generalized correlation function is called *correntropy*. Under this pair-wise independent condition, the centered correntropy reduces to zero since it is defined as the difference of correntropy and the information potential. In the conventional second order random processes, only pair-wise uncorrelatedness is required to zero the covariance function. The condition of pair-wise independence is much stronger than the pair-wise uncorrelatedness, this shows that the centered correntropy requires high order statistical information in order to attain zero. This can also been seen from the following observation.

Applying Taylor series expansion to the Gaussian kernel, we can rewrite the correntropy function as

\[
V(t, s) = \frac{1}{\sqrt{2\pi\sigma}} \sum_{k=0}^{\infty} \frac{(-1)^k}{(2\sigma^2)^k k!} E[(x_t - x_s)^{2k}] 
\]

which contains all the even-order moments of the random variable \( (x_t - x_s) \). Obviously different kernel functions would yield different expansions. But all the kernel functions mentioned above involve high order statistical information about random processes. Therefore the correntropy and centered correntropy partially characterizes the high order statistics of random processes. The correntropy and centered correntropy satisfy the following properties.

**Property 1:** The correntropy is positive definite and the centered correntropy is positive semi-definite.
Given a positive definite kernel function $\kappa(\cdot, \cdot)$, for any positive integer $n$, any points $t_1, \ldots, t_n$ in $\mathbb{T}$, and any not all zero real numbers $\alpha_1, \ldots, \alpha_n$, by definition we have

$$\sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \alpha_j \kappa(x_{t_i}, x_{t_j}) > 0.$$ 

Certainly, the expectation of any positive definite function is always positive definite. Thus we have

$$E \left[ \sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \alpha_j \kappa(x_{t_i}, x_{t_j}) \right] > 0.$$ 

This equals to

$$\sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \alpha_j E \left[ \kappa(x_{t_i}, x_{t_j}) \right] = \sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \alpha_j V(t_i, t_j) > 0. \quad (3\text{-}12)$$

Therefore the correntropy is positive definite. Similarly for the centered correntropy,

\[
0 \leq E \left[ \left\| \sum_{i=1}^{n} \alpha_i (\Phi(x_{t_i}) - E[\Phi(x_{t_i})]) \right\|^2 \right] \\
= E \left[ \left\langle \sum_{i=1}^{n} \alpha_i (\Phi(x_{t_i}) - E[\Phi(x_{t_i})]), \sum_{j=1}^{n} \alpha_j (\Phi(x_{t_j}) - E[\Phi(x_{t_j})]) \right\rangle \right] \\
= \sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \alpha_j E \left[ \left\langle \Phi(x_{t_i}) - E[\Phi(x_{t_i})], \Phi(x_{t_j}) - E[\Phi(x_{t_j})] \right\rangle \right] \\
= \sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \alpha_j U(t_i, t_j). \quad (3\text{-}13)
\]

Hence the centered correntropy is positive semi-definite.

The positive definiteness properties of the correntropy and centered correntropy are the most fundamental properties for these two functions because all the other following properties can be deduced from positive definiteness and more importantly it guarantees that the correntropy and centered correntropy uniquely determine two reproducing kernel Hilbert spaces. This positive definiteness property enables these two generalized functions a wide range of potential applications in statistical signal processing.

**Property 2:** $V(t, s)$ and $U(t, s)$ are symmetric: $V(t, s) = V(s, t)$, and $U(t, s) = U(s, t)$.
This is the direct consequence of symmetric kernel function used in the definitions of the correntropy and centered correntropy functions.

**Property 3:** \( V(t, t) > 0 \) and \( U(t, t) \geq 0 \).

Since \( \kappa(x_t, x_t) > 0 \) by the positive definiteness of kernel function, accordingly, \( V(t, t) > 0 \). For \( U(t, t) \), let \( \alpha_i, \alpha_j = 1 \) and \( n = 1 \) in Eq. (3–13), the result follows.

**Property 4:** \(|V(t, s)| < \sqrt{V(t, t)V(s, s)}\) and \(|U(t, s)| \leq \sqrt{U(t, t)U(s, s)}\).

Let \( n = 2 \) in Eq. (3–12) and Eq. (3–13), two expressions reduce to

\[
\alpha_1^2 V(t, t) + \alpha_2^2 V(s, s) > 2\alpha_1\alpha_2|V(t, s)|. \tag{3–14}
\]

\[
\beta_1^2 U(t, t) + \beta_2^2 U(s, s) \geq 2\beta_1\beta_2|U(t, s)|. \tag{3–15}
\]

We can substitute

\[
\alpha_1^2 = \frac{V(s, s)}{2\sqrt{V(t, t)V(s, s)}} \quad \text{and} \quad \alpha_2^2 = \frac{V(t, t)}{2\sqrt{V(t, t)V(s, s)}}
\]

\[
\beta_1^2 = \frac{U(s, s)}{2\sqrt{U(t, t)U(s, s)}} \quad \text{and} \quad \beta_2^2 = \frac{U(t, t)}{2\sqrt{U(t, t)U(s, s)}}
\]

into Eq. (3–14) and Eq. (3–15) respectively to obtain the properties above. ■

These properties are very similar to those of conventional correlation and covariance functions, but the correntropy and centered correntropy functions partially provide the high order statistics of random processes while the conventional correlation and covariance functions can only offer second order statistical information.

A random process \( \{x_t : t \in \mathbb{T}\} \) is said to be strictly stationary if the finite-dimensional joint probability density function is shift-invariant for each \( n \) and each choice of \( t_1, \ldots, t_n \) in \( \mathbb{T} \),

\[
f_n(x_1, \ldots, x_n; t_1, \ldots, t_n) = f_n(x_1, \ldots, x_n; t_1 + t_0, \ldots, t_n + t_0) \tag{3–16}
\]
for all $t_0$ in $T$. For a strict stationary random process, the correntropy and centered correntropy functions satisfy the following properties,

$$E[\Phi(x_t)] = \mu_{\Phi(x)}(0),$$  \hspace{1cm} (3–17)

$$V(t + \tau, t) = V(\tau, 0),$$  \hspace{1cm} (3–18)

and

$$U(t + \tau, t) = U(\tau, 0)$$ \hspace{1cm} (3–19)

for all $t$ and $\tau$ in $T$. We denote $\mu_{\Phi(x)}(0), V(t + \tau, t)$ and $U(t + \tau, t)$ by $\mu_{\Phi(x)}, V(\tau)$ and $U(\tau)$ respectively whenever Eq. (3–17) - Eq. (3–19) hold for all $t$ and $\tau$. It might be that the functions $\mu_{\Phi(x)}, V(\tau)$ and $U(\tau)$ are shift invariant in the sense of Eq. (3–17) - Eq. (3–19), yet the random process $\{x_t : t \in T\}$ is not strictly stationary. Since those equations represent properties of the random process that are of interest in their own right, we can define two forms of stationarity that are, in general, much weaker than the strict stationarity defined in Eq. (3–16), but much stronger than the conventional wide-sense stationarity and covariance stationarity since all the even-order moments should be time shift invariant from Eq. (3–11) in order to obtain a univariate correntropy function.

**Definitions:** A random process is said to be *correntropy-sense stationary (CSS)* if Eq. (3–17) and Eq. (3–18) hold for all $t$ and $\tau$; the process is said to be *centered correntropy stationary* if Eq. (3–19) holds for all $t$ and $\tau$.

Suppose $t = s + \tau$ in Eq. (3–6), then the centered correntropy becomes

$$U(s + \tau, s) = V(s + \tau, s) - \langle E[\Phi(x_{s+\tau})], E[\Phi(x_s)] \rangle.$$ \hspace{1cm} (3–20)

The right-hand side does not depend on $s$ for a correntropy-sense stationary process. Consequently, $U(s + \tau, s)$ does not depend on $s$. Therefore we have the following important fact:

*A correntropy-sense stationary process is also centered correntropy stationary.*
However, a centered correntropy stationary process might not be necessary correntropy-sense stationary. For a correntropy-sense stationary random process, the properties 2, 3 and 4 become

\[ V(\tau) = V(-\tau), \quad (3-21) \]
\[ V(0) > 0, \quad (3-22) \]
\[ |V(\tau)| < V(0), \quad (3-23) \]

for all \( \tau \).

The expectations, or ensemble averages, of a random process are average across the process. In practice, we do not have infinite realizations of random processes. Accordingly, we might use time averages to approximate the ensemble averages. In order to be rigorous for using this approach, we have to show that time averages converge to corresponding ensemble averages in some statistical sense. Consider a correntropy-sense stationary discrete-time random process \( \{x_n : n \in \mathbb{Z}^+\} \). Though the mean of the nonlinearly transformed random process \( \{\Phi(x_n) : n \in \mathbb{Z}^+\} \) is not required in applications, it is of interest to investigate the relationship between ensemble average and time average to the mean. In this regard, denote the mean of the process \( \{\Phi(x_n) : n \in \mathbb{Z}^+\} \), \( E[\Phi(x_n)] \) by \( \mu \) and define the time average to the mean as

\[ \hat{\mu}(N) = \frac{1}{N} \sum_{n=0}^{N-1} \Phi(x_n), \quad (3-24) \]

where \( N \) is the number of available samples used in the estimation. The estimator Eq. (3–24) is an unbiased estimator of the ensemble average of the process since

\[ E[\hat{\mu}(N)] = \mu \quad \text{for all } N. \quad (3-25) \]

Furthermore, the process is said to be mean-norm ergodic in the mean-norm-square error sense if the mean-norm-square value of the error between the ensemble average \( \mu \) and the
time average $\hat{\mu}(N)$ approaches zeros as the number of samples $N$ tends to infinity,

$$\lim_{N \to \infty} E \left[ \|\mu - \hat{\mu}(N)\|^2 \right] = 0. \quad (3\text{-}26)$$

Substituting Eq. (3–24) into Eq. (3–26), we can rewrite

\begin{align*}
E \left[ \|\mu - \hat{\mu}(N)\|^2 \right] &= E \left[ \left\| \mu - \frac{1}{N} \sum_{n=0}^{N-1} \Phi(x_n) \right\|^2 \right] \\
&= \frac{1}{N^2} E \left[ \left\| \sum_{n=0}^{N-1} (\Phi(x_n) - \mu) \right\|^2 \right] \\
&= \frac{1}{N^2} E \left[ \left\langle \sum_{n=0}^{N-1} (\Phi(x_n) - \mu), \sum_{k=0}^{N-1} (\Phi(x_k) - \mu) \right\rangle \right] \\
&= \frac{1}{N^2} \sum_{n=0}^{N-1} \sum_{k=0}^{N-1} E \left[ \langle \Phi(x_n) - \mu, \Phi(x_k) - \mu \rangle \right] \\
&= \frac{1}{N^2} \sum_{n=0}^{N-1} \sum_{k=0}^{N-1} U(n - k), \quad (3\text{-}27)
\end{align*}

where $U(n - k)$ is the centered correntropy at $n - k$ time lag for the correntropy-sense random process. Let $m = n - k$, the double summation in Eq. (3–27) can be simplified as follows,

$$E \left[ ||\mu - \hat{\mu}(N)||^2 \right] = \frac{1}{N} \sum_{\tau=-N+1}^{N-1} \left( 1 - \frac{|m|}{N} \right) U(m). \quad (3\text{-}28)$$

Hence, the necessary and sufficient condition for the process \{ $x_n : n \in \mathbb{Z}^+$ \} to be mean-norm ergodic in the mean-norm-square-error sense is that

$$\lim_{N \to \infty} \frac{1}{N} \sum_{\tau=-N+1}^{N-1} \left( 1 - \frac{\tau}{N} \right) U(m) = 0. \quad (3\text{-}29)$$

The time average $\hat{\mu}(N)$ of the process converges to the ensemble average $\mu$ in the mean-norm-square-error sense if the process \{ $x_n : n \in \mathbb{Z}^+$ \} is asymptotically correntropy-sense uncorrelated in the sense of Eq. (3–29).
The correntropy and centered correntropy can be estimated using time average for a discrete-time correntropy-sense stationary random process as follows:

\[
\hat{V}(l) = \frac{1}{N-l+1} \sum_{n=l}^{N-1} \kappa(x_n, x_{n-l}),
\]

and

\[
\hat{U}(l) = \frac{1}{N-l+1} \sum_{n=l}^{N-1} \kappa(x_n, x_{n-l}) - \frac{1}{N^2} \sum_{n=0}^{N-1} \sum_{m=0}^{N-1} \kappa(x_n, x_m)
\]

where \( \frac{1}{N^2} \sum_{n=0}^{N-1} \sum_{m=0}^{N-1} \kappa(x_n, x_m) \) is the estimate of information potential which is a constant depending on the signal and the specific kernel function used.

In what follows, we conduct several simulations to demonstrate some of the properties of correntropy and centered-correntropy. First, we would like to illustrate the relation between correntropy and centered-correntropy functions. 3 sets of 100000 i.i.d. data samples each of Gaussian, exponential, and Gamma distributed respectively were generated. The Gaussian source is set to be zero mean and unit variance. The parameter in exponential distribution is set to be 20. The parameters in Gamma distribution are set to be 0.2 and 1 respectively. Then we pass these data through an IIR filter with transfer function

\[
H(Z) = \frac{1 + 0.2Z^{-1}}{1 - 1.5Z^{-1} + 0.8Z^{-2}}.
\]

The i.i.d. signal and filtered signal are both normalized to zero mean and unit variance. A Gaussian kernel with unit kernel width is used to estimate the correntropy and centered correntropy. In Fig. 3-1, we plot the correntropy \([65]\) and centered correntropy functions for the original and filtered signal respectively. As proved in Eq. (3–10), under the pair-wise independent assumption, the value of correntorpy at non-zero lag is the information potential and the centered correntropy at non-zero lag reduce to zero since it is defined as the difference between correntropy and information potential. The left plot clearly illustrates the point. The estimated information potentials are 0.22803, 0.26098 and 0.31452 for Gaussian, exponential and Gamma distributed signals respectively. Likewise,
Figure 3-1. Correntropy and centered correntropy for i.i.d. and filtered signals versus the time lag

The right plot shows the correntropy and centered correntropy for the filtered signals. We will only plot the correntropy function in the following simulations since the difference between correntropy and centered correntropy is only the information potential which is a constant depending on the kernel and signal (Eq. (1–31)).

Our second simulation demonstrates the effectiveness of correntropy and centered correntropy in partial characterization of high order statistics of random processes. We compare the conventional autocorrelation and correntropy functions for two data sets. The first data set is 100000 i.i.d. data of Gaussian distributed with zero mean and unit variance. The second data set is the ARCH (autoregressive conditional heteroskedasticity) model which is used in econometrics to predict asset return volatility [77]. The ARCH time series is uncorrelated (second order statistics) but not i.i.d.. The time series model is
Figure 3-2. Autocorrelation and correntropy for i.i.d. and ARCH series versus the time lag.

defined as

\[ z_t = e_t \sqrt{\alpha_0 + \alpha_1 z_{t-1}^2 + \ldots + \alpha_r z_{t-r}^2}, \]  

(3–33)

where \( \{e_t\}_{t=0}^{\infty} \) is a white noise stochastic process and \( \alpha_0 > 0, \alpha_i \geq 0 \ \forall \ i = 1, \ldots, r \). We choose \( \{e_t\}_{t=0}^{\infty} \) to be i.i.d. Gaussian distribution of zero mean and unit variance, \( r = 1 \), \( \alpha_0 = 0.2 \) and \( \alpha_1 = 1 \). A Gaussian kernel with unit kernel width is used to estimate the correntropy. In Fig. 3-2, the autocorrelation functions for the i.i.d. Gaussian data and uncorrelated but dependent ARCH time series are given in the top two plots. It is expected that the conventional autocorrelation functions for an i.i.d. signal and an uncorrelated signal are the same since autocorrelation function only specifies the second order statistics. The bottom two plots show the correntropy functions for the same i.i.d. and uncorrelated signals respectively. As has been pointed out previously, the correntropy
partially characterizes the high order statistics. Accordingly, unlike the conventional correlation function, the correntropy would be different for i.i.d. and uncorrelated signals. We define a significance region to quantify the significant non-zero lag values. The maximum value of the significance region is the correntropy values at zero lag, which is 0.3989 in this specific simulation; the minimum value of the region is the 10 percent of difference between zero lag correntropy and estimate of information potential above the estimate of information potential, which is 0.3140. If any non-zero lag correntropy value falls into this region, we call it 90% significance which indicates there exists a “correlation” at that time lag. In this simulation, the correntropy values at time lag 1 and 2 are 0.3232 and 0.3156 respectively which fall into the significance region. This result clearly suggests that there exist a “generalized correlation” for this ARCH time series using correntropy quantification. Therefore correntropy specifies more information than second-order statistics compared to the conventional autocorrelation function.

The third simulation investigates the effectiveness of correntropy in capturing the PDF and nonlinearity information of the random processes. We generate 3 sets of 100000 i.i.d. data samples each of Gaussian, exponential, and Gamma distribution respectively to test correntropy’s ability to characterize PDF information of random processes. The Gaussian source is set to be zero mean and unit variance. The parameter in exponential distribution is set to be 20. The parameters in Gamma distribution are set to be 0.2 and 1 respectively. Then the data set is passed through the IIR filter Eq. (3–32). A Gaussian kernel with unit kernel width is used to estimate the correntropy. In Fig. 3-3 (a-d), we plot the conventional autocorrelation function and correntropy function for the original and filtered signal respectively. Because the conventional autocorrelation function only represents the time structure of random processes while containing no information about the PDF of random variables. Consequently, the autocorrelation functions for random processes of different PDF distributions are the same. This is demonstrated in the plots (a) and (c) for i.i.d. data of different distributions and filtered signals. However,
Figure 3-3. Autocorrelation and correntropy for i.i.d. and linearly filtered signals and Lorenz dynamic system versus the time lag.
the correntropy function not only captures the time structure of random processes, but also depicts the PDF distributions of random variables. The differences among the random processes of different PDF distributions can be easily seen in the plot (b). The correntropy values at non-zero lags are the estimate of information potential for different distributions Eq. (1–31), which are certainly different for different PDF distributions. Correntropy also contains the time structure of signals. Comparing (c) and (d) plots, we can see that the shape of correntropy is very similar to that of autocorrelation function. Moreover, correntropy functions are different for 3 different sources. The reason that the separation of correntropy functions for 3 different signals are not as obvious as the one in i.i.d. case (plot (b)) is because the distributions for the 3 filtered signals become more similar. In the plots (e) and (f), we demonstrate the difference of conventional correlation function and correntropy in capturing the nonlinearity intrinsic to systems. The Lorenz dynamic system is used here for illustration. The system function of Lorenz time series is given by

\[
\begin{align*}
\dot{x} &= \sigma (y - x), \\
\dot{y} &= -y - xz + Rx, \\
\dot{z} &= xy - bz
\end{align*}
\]

(3–34)

where \( R = 28, \sigma = 10 \) and \( b = 8/3 \). This set of parameter makes Lorenz dynamic system exhibit chaotic behavior. 10000 samples of the Lorenz dynamic system is generated by solving the equations with 4th-order Runge-Kutta method with integral step 0.01. Afterwards, it is normalized to zero mean and unit variance. In Fig. 3-3 (e-f), we plot the conventional autocorrelation function and correntropy for the 3 components of the dynamic system. Observe that the correntropy functions for the 3 state variables follow the same trend. They peak and drop around the same time lag. From the dynamic equations, it shows that \( x \) is nonlinearly coupled into \( y \) and \( z \). The periodic similarities in one state variable affect the other states. This is clearly demonstrated in the correntropy
functions. However, the conventional autocorrelation functions do not exhibit any periodic similarities among the three states. This simulation suggests that correntropy is able to extract the nonlinear coupling information embedded in the time structure of the time series while the conventional autocorrelation fails.

In the previous simulations, we just used Gaussian kernel with same kernel width. In fact, the kernel functions and kernel width play a crucial role in the correntropy and centered correntropy functions. Obviously, different kernel widths result in different estimate of information potential. Moreover, kernel width controls the ability of correntropy to capture the nonlinearity embedded in systems. From the RKHS perspective, different kernel functions determine different reproducing kernel Hilbert spaces and the kernel width defines the norm and inner product of RKHS. We will explore more on this issue in the next chapter. Here we first present an experiment on the different kernel width for the Gaussian kernel function. The same data sets on linear i.i.d. and the Lorenz time series from the previous example are used here. In Fig. 3-4, we plot the correntropy functions for i.i.d. and Lorenz time series using Gaussian kernel with different kernel width ranging from 0.01 to 15. It can be seen from the top two plots that correntropy loses the ability to differentiate time series of different PDF distributions and to detect the nonlinearity embedded in Lorenz time series when kernel width is too small ($\sigma = 0.01$ and 0.1 in this experiment). If the kernel is too big ($\sigma = 5$ and 15 in this experiment), the correntropy functions approach to conventional autocorrelation. The shape of correntropy in the bottom two plots are very similar to the ones of Fig. 3-3 (a) and (e). This can also be verified by the Taylor series expansion for the correntropy using Gaussian kernel in Eq. Eq. (3–11). If the kernel width is too large, the values of high order terms decay rapidly. Therefore correntropy approaches to conventional autocorrelation. The middle two plots give the correntropy functions of appropriate kernel width. It can be seen that correntropy can nicely separate the i.i.d signal of different PDF distributions and detect the embedded nonlinear coupling in the Lorenz time series.
To further quantify the relationship between kernel width and the ability of correntropy in separation i.i.d. signals of different PDF distributions, we define a separation coefficient $S(\sigma)$ as a function of kernel width $\sigma$ among the signals of different PDF distributions by

$$S(\sigma) = \sum_{i=1}^{L} \sum_{j>i}^{L} \frac{|\hat{I}_\sigma(i) - \hat{I}_\sigma(j)|}{|V_\sigma(0) - \min_i \hat{I}_\sigma(i)|},$$

where $L$ is number of i.i.d. signals, $\hat{I}_\sigma(i)$ is the estimate of information potential of $i^{th}$ signal (Eq. (1–31)), and $V_\sigma(0)$ is the correntropy at zero lag. The correntropy values at zero lag for different i.i.d. signals are the same since Gaussian kernel is isotropic. The separation coefficient basically computes the sum of normalized distance between different i.i.d. signals. Fig. 3-5 plots the separation coefficient as a function of kernel width $\sigma$. It can be easily seen that the best separation coefficient achieves at $\sigma = 1$. 
Next, we investigate the effect of different kernel functions used in the correntropy functions. The same data sets are used again, and we compare four different kernel functions, Gaussian, polynomial, sigmoid and . For a complete description, six kernel function expressions are given here.

1. Gaussian kernel : $\kappa(x_t, x_s) = \frac{1}{\sqrt{2\pi\sigma}} \exp\left\{ -\frac{(x_t - x_s)^2}{2\sigma^2} \right\}$ \hspace{1cm} (3–36)

2. Polynomial kernel : $\kappa(x_t, x_s) = (1 + x_t \cdot x_s)^d$ \hspace{1cm} (3–37)

3. Sigmoid kernel : $\kappa(x_t, x_s) = \tanh(\beta_0 x_t \cdot x_s + \beta_1)$ \hspace{1cm} (3–38)

4. Wave kernel : $\kappa(x_t, x_s) = \frac{1}{\alpha} \text{sinc} \left( \frac{x_t - x_s}{\alpha} \right)$ \hspace{1cm} (3–39)

5. Exponential kernel : $\kappa(x_t, x_s) = \lambda \exp\left( -\lambda|x_t - x_s| \right)$ \hspace{1cm} (3–40)

6. Inverse multiquadrics kernel : $\kappa(x_t, x_s) = \frac{1}{\sqrt{\theta^2 + (x_t - x_s)^2}}$ \hspace{1cm} (3–41)

In order to have a fair comparison, we choose suitable parameter for each of kernel functions for this specific data set. We select $\sigma = 1$ for Gaussian kernel, $d = 2$ for polynomial kernel, $\beta_0 = 10$ and $\beta_1 = 0.5$ for sigmoid kernel, $\alpha = 2.5$ for wave kernel,
Figure 3-6. Correntropy for i.i.d. signal and Lorenz time series with different kernel functions
\[ \lambda = 2.5 \text{ for exponential kernel and } \theta = 0.7 \text{ for inverse multiquadrics kernel.} \]

The correntropy functions for i.i.d. data of Gaussian, exponential and Gamma distributions and Lorenz time series are plotted using different kernel function in Fig. 3-6. Notice that all the kernel functions except polynomial kernel can separate different PDF data. The sigmoid kernel fails to detect the nonlinear coupling embedded in Lorenz time series. The wave, exponential and inverse multiquadrics kernels exhibit reasonable performance compared to Gaussian kernel.

### 3.2 Frequency-Domain Analysis

We have already presented the definitions and properties of correntropy and centered correntropy functions in the previous sections. If we view the correntropy and centered correntropy functions as generalized correlation and covariance functions, certainly we can proceed to the frequency-domain analysis. Similar to the conventional (power) spectral density function defined for a wide-sense stationary random processes, we can also define (correntropy) spectral density function for a correntropy-sense stationary random processes. Fourier transformation techniques offer an alternative approach to the time-domain analysis of correntropy for random processes.

**Definition:** Given a correntropy-sense stationary random process \( \{x_t, t \in \mathbb{T}\} \) with centered correntropy function \( U(\tau) \), the **correntropy spectral density function** is defined by

\[
P(\omega) = \int_{-\infty}^{\infty} U(\tau) e^{-j\omega \tau} d\tau,
\]

whenever the integral exists. In other word, the correntropy spectral density function is the Fourier transform of the centered correntropy function.

The variable \( \omega \) in Eq. (3–42) is usually called the *radian frequency* and is measured in radians per second. The frequency \( \omega/2\pi \), denoted as \( f \), is referred to as the “usual” frequency and measured in hertz(Hz). To convert an expression for the correntropy spectral density as a function of \( \omega \) into an expression for the same spectral density as a function of \( f \), one needs to replace \( \omega \) by \( 2\pi f \). Formally, the correntropy spectral density as
a function of frequency in Hz is defined by

\[ P(f) = \int_{-\infty}^{\infty} U(\tau)e^{-j2\pi f \tau} d\tau. \quad (3–43) \]

Since \( P(\omega) = \mathcal{F}\{U(\tau)\} \), we can obviously define the inverse transform of the correntropy spectral density function, \( U(\tau) = \mathcal{F}^{-1}\{P(\omega)\} \), as

\[
U(\tau) = \int_{-\infty}^{\infty} P(\omega)e^{j\omega\tau} d\omega/2\pi \\
= \int_{-\infty}^{\infty} P(f)e^{j2\pi f \tau} df. \quad (3–44)
\]

In particular, when the time lag \( \tau = 0 \), the correntropy spectral spectral density function becomes

\[
E[\kappa(x_t, x_t)] - E_{x_t}E_{x_t}[\kappa(x_t, x_t)] = E[\|\Phi(x_t)\|^2] - \|E[\Phi(x_t)]\|^2 = U(0) \\
= \frac{1}{2\pi} \int_{-\infty}^{\infty} P(\omega)e^0 d\omega \\
= \frac{1}{2\pi} \int_{-\infty}^{\infty} P(\omega) d\omega = \int_{-\infty}^{\infty} P(f) df. \quad (3–46)
\]

This equation shows that the difference between the expectation of norm square and norm square of the expectation of the nonlinearly transformed random process is the total area under the correntropy spectral density function \( P(f) \) of the process. In second-order random processes analysis, the power in the random process is the total area under the power spectral density function. By the similar approach, we define the difference between the expectation of norm square and norm square of the expectation of the nonlinearly transformed random process, in other word, the centered correntropy at zero lag \( U(0) \), as the correntropy power. From a geometrical perspective, \( U(0) \) can also be considered as “generalized” variance, or “generalized” power, of the nonlinearly transformed random process in the feature space. Physically, \( P(\omega) \) represents the density of correntropy power at frequency \( \omega \) radians/sec, and \( P(f) \) plays the same role for frequency in Hz. The correntropy power in any frequency band is obtained by integrating the correntropy
spectral density function over the range of frequencies that build up the band. Specifically, the correntropy power in a frequency band from $\omega_1$ to $\omega_2$ radians/sec, or from $f_1$ to $f_2$ Hz, is defined as

$$\frac{1}{2\pi} \int_{\omega_1}^{\omega_2} P(\omega) d\omega = \int_{f_1}^{f_2} P(f) df. \quad (3-47)$$

The correntropy spectral density function for a correntropy-sense stationary random process satisfies the following properties:

**Property 1:** $P(\omega) = P(-\omega)$, for all $\omega$.

Because $U(\tau) = U(-\tau)$ for a correntropy-sense stationary random process, we then have

$$P(\omega) = \int_{-\infty}^{\infty} U(\tau)e^{-j\omega\tau} d\tau = \int_{-\infty}^{\infty} U(-\lambda)e^{+j\omega\lambda} d\lambda$$

$$= \int_{-\infty}^{\infty} U(\lambda)e^{-j(-\omega)\lambda} d\lambda = P(-\omega). \quad (3-48)$$

**Property 2:** $P(\omega) = [P(\omega)]^*$, for all $\omega$.

$[P(\omega)]^*$ denotes the complex conjugate of $P(\omega)$. To prove this property, notice that

$$[P(\omega)]^* = \left[ \int_{-\infty}^{\infty} U(\tau)e^{-j\omega\tau} d\tau \right]^* = \int_{-\infty}^{\infty} U(\tau)[e^{-j\omega\tau}]^* d\tau$$

$$= \int_{-\infty}^{\infty} U(\tau)e^{+j\omega\tau} d\tau = \int_{-\infty}^{\infty} U(\tau)e^{-j(-\omega)\tau} d\tau$$

$$= P(-\omega) = P(\omega). \quad (3-49)$$

This property shows that the correntropy spectral density function is real.

**Property 3:** $P(\omega) \geq 0$, for all $\omega$. 
Observe that for any $T \geq 0$,

$$0 \leq E\left\{\left\| \int_0^T (\Phi(x_t) - E[\Phi(x_t)]) e^{-j\omega t} dt \right\|^2 \right\}$$

$$= E\left\{\left[ \int_0^T (\Phi(x_t) - E[\Phi(x_t)]) e^{-j\omega t} dt \right], \left[ \int_0^T (\Phi(x_s) - E[\Phi(x_s)]) e^{-j\omega s} ds \right]^* \right\}$$

$$= E\left\{ \int_0^T \int_0^T \langle \Phi(x_t) - E[\Phi(x_t)], \Phi(x_s) - E[\Phi(x_s)] \rangle e^{-j\omega(t-s)} dt ds \right\}$$

$$= \int_0^T \int_0^T U(t-s)e^{-j\omega(t-s)} dt ds.$$ 

Hence, this last double integral is nonnegative. Since $T \geq 0$, certainly, $T^{-1}$ is also greater than 0, and this results in

$$T^{-1} \int_0^T \int_0^T U(t-s)e^{-j\omega(t-s)} dt ds \geq 0 \quad (3\text{-}50)$$

for all $T \geq 0$. We make a change-of-variable transformation in Eq. (3-50) by letting $t - s = \tau$. The transformation maps the original region of integration $\{(t, s) : 0 \leq t \leq T, 0 \leq s \leq T\}$ onto the region $\{(t, \tau) : 0 \leq t \leq T, t - T \leq \tau \leq t\}$. In other word, the original region of square is changed into a parallelogram, which can also be decomposed into two regions with $G_1 = \{(t, \tau) : \tau \leq t \leq T, 0 \leq \tau \leq T\}$ and $G_2 = \{(t, \tau) : 0 \leq t \leq T + \tau, -T \leq \tau \leq 0\}$. Then Eq. (3-50) can be rewritten as the sum of an integral over the region $G_1$ and an integral over the region $G_2$ as follows:

$$T^{-1} \int_0^T \int_0^T U(t-s)e^{-j\omega(t-s)} dt ds$$

$$= T^{-1}\left\{ \int_0^T \left[ \int_0^\tau U(\tau)e^{-j\omega \tau} d\tau \right] d\tau + \int_0^T \left[ \int_{-T}^{T+\tau} U(\tau)e^{-j\omega \tau} d\tau \right] d\tau \right\}$$

$$= T^{-1}\left\{ \int_0^T (T-\tau)U(\tau)e^{-j\omega \tau} d\tau + \int_0^T (T+\tau)U(\tau)e^{-j\omega \tau} d\tau \right\}$$

$$= \int_{-T}^T \frac{(T - |\tau|)}{T} U(\tau)e^{-j\omega \tau} d\tau. \quad (3\text{-}51)$$
Let us define the function $\hat{U}_T(\tau)$ as

$$
\hat{U}_T(\tau) = \begin{cases} 
1 - \frac{|\tau|}{T}, & |\tau| \leq T \\
0, & |\tau| > T 
\end{cases}
$$

(3–52)

Then for any fixed $\tau$, we have

$$
\lim_{T \to \infty} \hat{U}_T(\tau) = U(\tau).
$$

(3–53)

Therefore, the limit of Eq. (3–51) becomes

$$
\lim_{T \to \infty} \int_{-T}^{T} \frac{(T - |\tau|)}{T} U(\tau)e^{-j\omega \tau} d\tau = \lim_{T \to \infty} \int_{-\infty}^{\infty} \hat{U}_T(\tau)e^{-j\omega \tau} d\tau = \int_{-\infty}^{\infty} \lim_{T \to \infty} \hat{U}_T(\tau)e^{-j\omega \tau} d\tau = \int_{-\infty}^{\infty} U(\tau)e^{-j\omega \tau} d\tau = P(\omega).
$$

(3–54)

The interchange of order of the limit and the integration is validated by the dominated convergence theorem [21]. Eq. (3–51) and Eq. (3–54) established that $P(\omega)$ is the limit of a sequence of nonnegative numbers for any $\omega$, and the limit of a sequence of nonnegative numbers must be nonnegative. Consequently, $P(\omega) \geq 0$. 

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CHAPTER 4
CORRENTROPY ANALYSIS BASED ON RKHS APPROACH

From the definitions of correntropy and centered correntropy functions, we know that there are two reproducing kernel Hilbert spaces embedded in the correntropy and centered correntropy functions. The first RKHS is induced by the symmetric positive definite data independent kernel functions, like the Gaussian kernel, Sigmoid kernel and others Eq. (3–36) - Eq. (3–41). These kernel functions nonlinearly transform the original random processes into a high dimensional RKHS, which is called feature space. The linear operations on the transformed random processes will become nonlinear operations on the original random processes when mapped back by the inverse nonlinear transformation.

The second RKHS is induced by the symmetric positive definite data dependent correntropy and centered correntropy functions directly. Unlike the conventional autocorrelation and covariance functions that operate directly on the original random processes and determine data dependent RKHS, correntropy and centered correntropy operate on the expectation of nonlinearly transformed random processes, hence induce reproducing kernel Hilbert spaces that are very different from the ones determined by data independent kernel functions and conventional autocorrelation and covariance functions. The inclusion of data independent kernel functions inside the expectation operation makes the correntropy and centered correntropy functions different from kernel functions and expectation operation alone. This also changes the dynamics of the RKHS induced by the correntropy and centered correntropy and makes it unique.

In this chapter, we investigate the correntropy and centered correntropy functions from a geometrical perspective by analyzing the reproducing kernel Hilbert spaces induced by them directly and the one by kernel functions alone. These two approaches to analyze the correntropy and centered correntropy will lead us to a better understanding of the geometrical structure of the RKHS induced by kernel functions, correntropy and centered correntropy respectively.
By the Mercer’s theorem, any symmetric positive definite functions possess an
eigen-decomposition. Even without the explicit knowledge of the eigen-functions and
eigen-values, kernel methods solve problems which can be expressed in terms of inner
product because the evaluation of the inner product is equivalent to the value of kernel
function. In this chapter, we present a new method to explicitly build the RKHS with
Gaussian kernel based on polynomial functions. A knowledge of the bases that construct
the RKHS offer a potential applications of RKHS beyond the inner product.

4.1 RKHS Induced by the Kernel Function

By the Mercer’s theorem in chapter 1 Eq. (1–14), any symmetric positive definite
function $\kappa(x_t, x_s)$ can be rewritten as an inner product between two vectors in the feature
space, i.e.,

$$
\kappa(x_t, x_s) = \sum_{k=0}^{\infty} \lambda_k \varphi_k(x_t) \varphi_k(x_s)
= \langle \Phi(x_t), \Phi(x_s) \rangle
$$

(4–1)

\[ \Phi : x_t \mapsto \sqrt{\lambda_k} \varphi_k(x_t), \quad k = 1, 2, \ldots, \text{and } t \in \mathbb{T}. \]

where $\lambda_k$ and $\varphi_k$ are eigen-values and orthogonal eigen-functions for the kernel $\kappa(\cdot, \cdot)$
respectively. The nonlinear transformation $\Phi$ maps the random process \{x_t : t \in \mathbb{T}\} into
another random process \{\Phi(x_t) : t \in \mathbb{T}\} in the high-dimensional feature space, which is a
reproducing kernel Hilbert space induced by the kernel according to the Moore-Aronzajn
theorem Eq. (1–2). The construction of RKHS based upon the eigen-values and
eigen-functions $\lambda_k$ and $\varphi_k$ follows the same approach from Eq. (1–14) to Eq. (1–20)
by substituting the autocorrelation function $R(t, s)$ with kernel function $\kappa(x_t, x_s)$. Notice
that if isotropic kernel functions are used, specifically the Gaussian kernel Eq. (3–36),
wave kernel Eq. (3–39), exponential kernel Eq. (3–40) and inverse multiquadrics kernel
Eq. (3–41), then the norm of transformed random process $\|\Phi(x_t)\|^2$ is a constant equals
to $\kappa(0)$. This shows that the transformed random process \{\Phi(x_t) : t \in \mathbb{T}\} resides
on the sphere. The isotropic kernel functions transform the random processes such
that the instant power become constant. This approach offers potential applications in communications and signal processing.

The data independent kernel functions include Gaussian kernel, Sigmoid kernel and others Eq. (3–36) - Eq. (3–42). These data independent kernel functions are embedded inside the correntropy and centered correntropy functions. An analysis of correntropy and centered correntropy from kernel perspective offers a new insight.

4.1.1 Correntropy Revisited from Kernel Perspective

According to the inner product expression in Eq. (4–1), we can rewrite the correntropy in terms of the nonlinear mapping \( \Phi \) as

\[
V(t, s) = E[\langle \Phi(x_t), \Phi(x_s) \rangle].
\] (4–2)

Likewise the centered correntropy can also be expressed as

\[
U(t, s) = E[\langle \Phi(x_t), \Phi(x_s) \rangle] - \langle E[\Phi(x_t)], E[\Phi(x_s)] \rangle
= E[\langle \Phi(x_t) - E[\Phi(x_t)], \Phi(x_s) - E[\Phi(x_s)] \rangle]
\] (4–3)

It can be seen that the correntropy function is a “conventional” correlation function for the transformed random process in the high-dimensional RKHS. While the centered correntropy is nothing but the correntropy for the zero mean (centered) random process \( \{ \Phi(x_t) - E[\Phi(x_t)] : t \in \mathbb{T} \} \). This way of defining the generalized correlation and covariance functions follows the same spirit of the standard correlation and covariance functions. The high order statistics for any random process in the input space turns out to be the “second-order” statistics in the feature space. Kernel-based learning algorithms employ the nonlinear mapping \( \Phi \) to treat nonlinear algorithms in a linear way if the problems can be expressed in terms of inner product [49]. This suggests that we can deal with nonlinear systems efficiently and elegantly in a linear fashion when applying the correntropy and centered correntropy functions.
In fact, all the previous properties of correntropy and centered correntropy functions in section 2.1 can be derived in a kernel framework. For example, property 3 can be shown that \( V(t, t) = E[\|\Phi(x_t)\|^2] \) which means that \( V(t, t) \) is nothing but the expectation of norm square of transformed random process. The centered correntropy at zero lag \( U(t, t) \) can be rewritten as

\[
U(t, t) = E \left[ \langle \Phi(x_t) - E[\Phi(x_t)], \Phi(x_t) - E[\Phi(x_t)] \rangle \right]
\]
\[
= E \left[ \| \Phi(x_t) - E[\Phi(x_t)] \|^2 \right]
\]
\[
= E[\|\Phi(x_t)\|^2] - \|E[\Phi(x_t)]\|^2
\] (4–4)

Therefore, the centered correntropy can be viewed as the expectation of the norm square of zero-mean (centered) transformed random process or the generalized covariance of transformed random process. Similarly, property 4 can be cast in terms of the nonlinearly transformed random process. The inequality can be re-expressed as

\[
E[\langle \Phi(x_t), \Phi(x_s) \rangle] < \sqrt{E[\|\Phi(x_t)\|^2]E[\|\Phi(x_s)\|^2]},
\]

which is called the *generalized Cauchy-Schwarz inequality* in the reproducing kernel Hilbert space.

From chapter 3.1, we know that if the random process is pair-wise independent then the correntropy at non-zero lags becomes the information potential and the centered correntropy at non-zero lags reduces to zero. Under this condition, the nonlinear transformed random process becomes “uncorrelated” in the feature space. The condition of pair-wise independence in the original input space implies pair-wise uncorrelatedness in the feature space, but not vice versa. This offers correntropy and centered correntropy a wide range of potential applications in machine learning and signal processing.

The kernel function perspective to correntropy and centered correntropy suggests that we can treat the nested kernel function inside the expectation operator as an implicit nonlinear transformation that maps the original random process into the RKHS induced
by the kernel function, where the transformed random process resides on the sphere if isotropic kernel functions are used. Then correntropy and centered correntropy functions are nothing but “generalized” autocorrelation and covariance functions of the nonlinearly transformed random process. This nonlinear transformation of random processes brings a range of advantages, as what have been demonstrated in chapter 2, but also poses challenges about operations on those transformed random processes since only the inner product of the transformed random processes in known without any knowledge of the individual nonlinear transformations. A method of explicit construction of the nonlinear functionals would alleviate the problem.

4.1.2 An Explicit Construction of a Gaussian RKHS

The Mercer’s theorem does not explicitly provide the basis \( \varphi \) that builds the RKHS. Rather it only offers the inner product of the eigen-functions because it is sufficient for the statistical learning problems which can be expressed in terms of inner products. One of the most fundamental issues in learning problems is the selection of the data representation. In kernel-based learning algorithms, this translates into the choice of the functionals, or the appropriate feature space RKHS. The reason is that the nonlinear mapping has a direct impact on the kernel and thus, on the solution of the given learning problems. Different kernels (polynomial, sigmoid, Gaussian) very likely will result in different performances. The functional form of RKHS is still of great interest to optimize performance and to get an insight into the appropriateness of the data representation. Ultimately, this will allow us to utilize the RKHS structure and expand the class of algorithms, beyond inner products, that can be developed in kernel space. The advantage of kernel-based learning algorithms becomes also a disadvantage. The general question of how to select the ideal kernel for a given problem remains an open issue. Recently, there have been attempts to explicitly construct an RKHS. A. Rakotomamonjy et al. proposed a method of building an RKHS and its associated kernels by means of frame theory [78]. Any vector in that RKHS can be represented by linear combination of the
frame elements. But a frame is not necessarily linear independent although it results in stable representation.

In this section, we take the polynomial space approach to construct explicitly an RKHS associated with one of the most popular kernels, the Gaussian kernel. By transforming a generalized Fock space [45] with a positive operator, we build an RKHS associated with Gaussian kernel. The functionals, are explicitly given by the polynomials. Unlike the Mercer’s theorem approach, these functional are not necessary orthonormal. More importantly, we can gain control over the dimension of the RKHS by means of selecting the polynomial degree. The simulation suggests that the effective dimension of RKHS with Gaussian kernel is relatively small.

The definitions of functionals and inner product of a general Hilbert space are given. Then, a kernel function is imposed on this general Hilbert space to make it a reproducing kernel Hilbert space. This approach of building an RKHS with polynomials can also be found in [45], which is called generalized Fock space. Our contribution is that it is an RKHS associated with Gaussian kernel that we explicitly construct by introducing new definitions of functionals and kernel function.

First we construct an inner product space $\mathcal{H}$ by defining functionals and inner product. The evaluation of functional $f$ at any given $x$ is given by

$$f(x) = e^{-\frac{x^2}{2\sigma_0}} \sum_{k=0}^{n} \frac{f_k}{k!} x^k,$$

where $\sigma_0$ is a constant and $(n + 1)$-tuple $(f_0, f_1, \ldots, f_n)$ are the coefficients which uniquely characterize the polynomial $f$. Then the inner product between any two functionals $f$ and $h$ can be specified in the form

$$\langle f, h \rangle = \sum_{k=0}^{n} \frac{\sigma_k}{k!} f_k h_k,$$
where $f_k$ and $h_k$ are coefficients for $f$ and $h$ respectively and $\sigma = (\sigma_0, \sigma_1, \ldots, \sigma_n)$ is a set of positive constants chosen \textit{a priori}. It can be easily seen that this inner product space $\mathcal{H}$ is complete thus forming a Hilbert space.

In order to make $\mathcal{H}$ a reproducing kernel Hilbert space, we impose a kernel function $\kappa$ on $\mathcal{H}$ in the following form

$$
\kappa(x, y) = e^{-\frac{x^2+y^2}{2\sigma_0}} \sum_{k=0}^{n} \frac{1}{k!\sigma_k} (xy)^k.
$$

(4–7)

It can be verified that the Hilbert space $\mathcal{H}$, equipped with such $\kappa$, is a \textit{reproducing kernel Hilbert space} and the kernel function $\kappa(x, \cdot)$ is a \textit{reproducing kernel} because of the following two properties of $\kappa(x, y)$:

1. $\kappa(x, y)$ as a function of $y$ belongs to $\mathcal{H}$ for any fixed $x$ because we can rewrite $\kappa(x, y)$ as

$$
\kappa(x, \cdot)(y) = e^{-\frac{y^2}{2\sigma_0}} \sum_{k=0}^{n} \frac{(x^k \cdot e^{-\frac{x^2}{2\sigma_0}})}{k!} y^k. 
$$

(4–8)

i.e., the constants $(x^k/\sigma_k \cdot e^{-\frac{x^2}{2\sigma_0}})$, $k = 0, 1, \ldots, n$ become the coefficients $f_k$, $k = 0, 1, \ldots, n$ in the definition of $f$, and thus

$$
\kappa(x, \cdot) \in \mathcal{H}. 
$$

(4–9)

2. Given any $f \in \mathcal{H}$, the inner product between reproducing kernel and $f$ yields the function itself,

$$
< \kappa(x, \cdot), f > = \sum_{k=0}^{n} \frac{\sigma_k}{k!} (x^k \cdot e^{-\frac{x^2}{2\sigma_0}}) \cdot f_k
$$

$$
= e^{-\frac{x^2}{2\sigma_0}} \sum_{k=0}^{n} \frac{f_k}{k!} x^k = f(x).
$$

(4–10)

This is so called \textit{reproducing property}.

The RKHS constructed above has the freedom to choose the degree of functionals, i.e., the dimension $n$ of the kernel space $\mathcal{H}$. The most interesting case is that we might
extend it to an infinite-dimensional RKHS provided that the norm of functional is finite as
\( n \to \infty \), i.e., given a sequence of positive weighting constants satisfying certain conditions
\( \sigma = (\sigma_0, \sigma_1, \ldots) \),
\[
\| f \|^2 = \langle f, f \rangle = \sum_{k=0}^{\infty} \frac{\sigma_k}{k!} f_k^2 < \infty. \tag{4-11}
\]
Then the functionals, inner product and reproducing kernel in \( \mathcal{H} \) will be defined by Eq. (4-5), Eq. (4-6) and Eq. (4-7) with \( n = \infty \).

In the special situation of weights
\[
\sigma_k = \sigma_0^k, \quad k = 1, 2, \ldots \tag{4-12}
\]
where \( \sigma_0 \) is a fixed positive constant, then the reproducing kernel Eq. (4-7) in the
infinite-dimensional RKHS becomes
\[
\kappa(x, y) = e^{-\frac{x^2 + y^2}{2\sigma_0}} \sum_{k=0}^{\infty} \frac{1}{k!} \left( \frac{xy}{\sigma_0} \right)^k
= e^{-\frac{x^2 + y^2}{2\sigma_0}} e^{\frac{xy}{\sigma_0}}
= e^{-\frac{(x-y)^2}{2\sigma_0}}, \tag{4-13}
\]
which is the Gaussian kernel used widely in machine learning, function approximation,
density estimation, support vector machine, and etc. The constant \( \sigma_0 \) turns out to be the
kernel width. It controls the norm length of those functionals in RKHS, i.e., the spread of
nonlinear mapped data sample in feature space.

Comparing this method with Mercer’s theorem, we notice that there are two major
differences between them.

1. First, we have given an explicit expression for the functionals in the RKHS
associated with Gaussian kernel in terms of polynomials while Mercer’s theorem
never does that. We can get the exact evaluations for those functionals at each point
in the RKHS. This enables us to know exactly the structure of the RKHS associated with the Gaussian kernel.

2. Second, the functionals we constructed above are not necessary an orthonormal basis, while the Mercer’s theorem is realized by orthonormalizing the RKHS. This perspective provides also a general alternative to build an RKHS from known functionals besides from Mercer’s theorem.

3. Third, we can have control of the dimension of the RKHS by means of selecting the polynomial degrees \( n \) in Eq. (4–7).

The method we constructed an RKHS enables us to have the explicit expression of the functional in RKHS associated with Gaussian kernel. Hence we can exactly know the nonlinear mapping \( \Phi \) used in the kernel-based learning algorithm and so operate directly with the transformed data to extend the algorithms beyond inner products. Furthermore, as we have the control of the dimension of the RKHS \( \mathcal{H} \), this might help the computational complexity issue in kernel-based learning algorithms through approximation of Gaussian kernel by polynomials as indicated in equation Eq. (4–7).

The previous way of explicitly constructing an RKHS by means of polynomial functions was employed by De Figueiredo [45] which is based on the Fock space. The idea of Fock space was first proposed by Fock in [79] to be used in quantum mechanics, where quantum states are described in the way of passing from one single object to collections of objects. More recently Figueiredo introduced an “arbitrarily weighted Fock space”, which was called generalized Fock space in [45]. The space is equipped with an appropriate weighted inner product, thus forming an RKHS. The proposed RKHS has been used in learn/nonlinear system and signal analysis, where a number of problems are involving approximation and inversion of nonlinear functions/functionals, and nonlinear operators. In the univariate case, a generalized Fock space \( F^n \) is an RKHS, where the functionals \( f \),
The inner product and kernel function $F$ are defined as follows respectively,

$$f(u) = \sum_{k=0}^{n} \frac{f_k}{k!} u^k,$$

(4–14)

$$\langle f, h \rangle_F = \sum_{k=0}^{n} \frac{\sigma_k}{k!} f_k h_k,$$

(4–15)

$$F(u, v) = \sum_{k=0}^{n} \frac{1}{k! \sigma_k} (uv)^k,$$

(4–16)

where the real $(n + 1)$-tuple $(f_0, f_1, \ldots, f_n)$ completely characterizes $f$, $\sigma = (\sigma_0, \sigma_1, \ldots, \sigma_n)$ is a set of positive weighting constants which are chosen \textit{a priori} according to the problems under consideration. It can be shown that this generalized Fock space is an RKHS. Similar to the RKHS $\mathcal{H}$ we constructed above, the generalized Fock space $F^n$ has the freedom of choosing the space dimension. The interesting case is that when the space becomes infinite dimensional while the norm of the functional satisfying the same condition Eq. (4–11) as $n \to \infty$. Then the kernel function $F(u, v)$, defined by Eq. (4–16), will become an exponential kernel as $n \to \infty$

$$\kappa(x, y) = e^{\frac{xy}{\sigma_0}},$$

(4–17)

given the same weights constraint as Eq. (4–12).

It can be noticed that there are similarity and difference between the RKHS $\mathcal{H}$ and the generalized Fock space $F^n$. The definitions for the inner product inside the two spaces are the same, while the functionals and kernel function are different. The relationship of the two spaces $\mathcal{H}$ and $F^n$ is connected by a theorem in [2], which states that if $H_1$ and $H_2$ are two reproducing kernel Hilbert spaces of the same definitions of inner product, then there exits a positive operator with bound not greater than 1 that transforms $H_1$ into $H_2 \subset H_1$. Comparing the definitions of functionals for two spaces, we can see that the $e^{-x^2/2\sigma_0}$ plays the role of a positive operator with bound not greater than 1, thus transforming the generalized Fock space $F^n$ into $\mathcal{H}$ such that $\mathcal{H} \subset F$. 
We present a simple simulation here to show the effectiveness of approximation of the polynomial functionals to the Gaussian kernel. In the simulation, we calculate the square error between the Gaussian kernel and the proposed kernel in Eq. (4–7) of order $n$. The kernel width is chosen to be 1, and the range of the calculated data is from -5 to 5. The Fig. 4-1 plots the square error versus order $n$. The line becomes flat at order 11 is due to the computation precision of MATLAB. The figure suggests that the effective dimension of the RKHS with Gaussian kernel is relatively small. With only order 11, we can effectively approximate the Gaussian kernel by polynomials. This also indicates that for practical purpose it is sufficient to work with much smaller dimensional space instead of infinite dimensional space for a Gaussian kernel in kernel-based learning algorithms.

4.2 RKHS Induced by Correntropy and Centered Correntropy Functions

We have seen that the kernel function nested inside the correntropy and centered correntropy functions uniquely induces a reproducing kernel Hilbert space. The nonlinear
transformation $\Phi$ maps the original random processes into the high-dimensional RKHS where the transformed random processes appear on a sphere if isotropic kernel function is used. The correntropy and centered correntropy can be carried out by operating on those transformed random processes. However, the correntropy and centered correntropy functions are symmetric non-negative definite themselves, hence determine two reproducing kernel Hilbert spaces as well. This perspective is in the same spirit of RKHS in statistical signal processing, where the autocorrelation and covariance functions are symmetric non-negative definite thus induce reproducing kernel Hilbert space (refer to the review in chapter 1.2). All the linear statistical signal processing using autocorrelation and covariance functions can be also treated as functional operations in RKHS. But the RKHS induced by correntropy and centered correntropy functions is different from the one induced by conventional autocorrelation and covariance functions. The RKHS by conventional autocorrelation and covariance functions is based on second order random processes, hence suitable for the Gaussian processes. While the RKHS by correntropy and centered correntropy function includes the high order statistical information of random processes, hence goes beyond the Gaussian processes. It is conjectured that RKHS by correntropy and centered correntropy functions might encompass a large class of processes. From the other perspective, the RKHS by correntropy and centered correntropy is also different from the one by kernel functions. One obvious characteristic is that RKHS by correntropy and centered correntropy is data dependent while the RKHS by kernel function is data independent. The inclusion of kernel function inside the expectation operator makes the correntropy and centered correntropy functions unique. It departs from the conventional RKHS induced by the autocorrelation and covariance functions used in statistical signal processing and also the data independent kernel functions used in kernel machine learning. In this section, we analyze the geometry of RKHS by correntropy and centered correntropy functions.
4.2.1 Geometry of Nonlinearly Transformed Random Processes

Given a random process \( \{ x_t : t \in T \} \), we can construct another nonlinearly transformed random process \( \{ \Phi(x_t) - E[\Phi(x_t)] : t \in T \} \), where \( \Phi \) is the nonlinear mapping associated with the kernel function Eq. (4–1). The reason to work with the zero-mean nonlinearly transformed random process shall become clear shortly. Let us define a linear manifold spanned by the random process \( \{ \Phi(x_t) - E[\Phi(x_t)] : t \in T \} \), denoted as \( L(\Phi(x_t) - E[\Phi(x_t)] : t \in T) \), to be the set of all random variables \( \xi \) that can be written as the form

\[
\xi = \sum_{i=1}^{n} c_i (\Phi(x_{t_i}) - E[\Phi(x_{t_i})]),
\]

(4–18)

for some integer \( n \), real constants \( c_1, \ldots, c_n \), and \( t_1, \ldots, t_n \in T \). In other words, \( L(\Phi(x_t) - E[\Phi(x_t)] : t \in T) \) contains all finite linear combinations of the random variables \( \{ \Phi(x_t) - E[\Phi(x_t)] : t \in T \} \). Close the set in Eq. (4–18) topologically according to the convergence in the mean using the norm

\[
\|\xi - \zeta\| = \sqrt{E[\|\xi - \zeta\|^2]}
\]

(4–19)

and denote the set of all linear combinations of random variables and its limit points by \( L_2(\Phi(x_t) - E[\Phi(x_t)] : t \in T) \). In other words, \( L_2(\Phi(x_t) - E[\Phi(x_t)] : t \in T) \) consists of all random variables in the linear manifold \( L(\Phi(x_t) - E[\Phi(x_t)] : t \in T) \), together with all random variables \( \xi \) such that there exists a sequence of random variables \( \xi_n \) in \( L(\Phi(x_t) - E[\Phi(x_t)] : t \in T) \) converging to \( \xi \), in the sense that \( \|\xi - \xi_n\| = \sqrt{E[\|\xi - \xi_n\|^2]} \to 0 \) as \( n \to 0 \).

It is well known from the theory of quadratically integrable functions that \( L_2(\Phi(x_t) - E[\Phi(x_t)] : t \in T) \) forms a Hilbert space if the inner product is the one corresponding to the norm in Eq. (4–19), namely

\[
\langle \xi, \zeta \rangle = E[\langle \xi, \zeta \rangle].
\]

(4–20)
Then by the definition of crosscorrentropy Eq. (3–8), we have

\[ \langle \xi, \zeta \rangle = E[\langle \xi, \zeta \rangle] = E[\langle \Phi(x) - E[\Phi(x)], \Phi(y) - E[\Phi(y)] \rangle] = E[\kappa(x, y)] - E_xE_y[\kappa(x, y)] = U(x, y). \quad (4–21) \]

In other words, the inner product in \( L_2(\Phi(x_t) - E[\Phi(x_t)] : t \in \mathbb{T}) \) is the centered correntropy function of the original random variables. This definition will bridge the space \( L_2(\Phi(x_t) - E[\Phi(x_t)] : t \in \mathbb{T}) \) and the RKHS induced by the centered correntropy function.

Now let us define a “generalized” deviation of any random variable \( \xi \) in \( L_2(\Phi(x_t) - E[\Phi(x_t)] : t \in \mathbb{T}) \) as

\[ \sigma(\xi) = \sqrt{E[\|\xi\|^2]} = \|\xi\| = \sqrt{E[\langle \Phi(x) - E[\Phi(x)], \Phi(x) - E[\Phi(x)] \rangle]} = \sqrt{E[\kappa(x, x)] - E_xE_y[\kappa(x, x)]} = \sqrt{U(x, x)}. \quad (4–22) \]
Consider the subspace $S$ spanned by two random variables $x_1$ and $x_2$ (we drop the time index here for simplicity without any ambiguity):

$$S = \{ c_1(\Phi(x_1) - E[\Phi(x_1)]) + c_2(\Phi(x_2) - E[\Phi(x_2)]) \}. \quad (4-23)$$

Then it makes sense to talk about the angle $\phi$ between two vectors $\xi_1$ and $\xi_2$ in $S$ (see Fig. 4-2). By plane trigonometry, we have

$$\sigma^2(\xi_1 - \xi_2) = \sigma^2(\xi_1) + \sigma^2(\xi_2) - 2\sigma(\xi_1)\sigma(\xi_2) \cos \phi, \quad (4-24)$$

which is the same as

$$\|\xi_1 - \xi_2\|^2 = \|\xi_1\|^2 + \|\xi_2\|^2 - 2\langle\xi_1, \xi_2\rangle = \|\xi_1\|^2 + \|\xi_2\|^2 - 2\|\xi_1\|\|\xi_2\| \cos \phi. \quad (4-25)$$

Consequently, we have

$$\cos \phi = \frac{\langle\xi_1, \xi_2\rangle}{\|\xi_1\|\|\xi_2\|} = \frac{U(x_1, x_2)}{\sqrt{U(x_1, x_1)U(x_2, x_2)}.} \quad (4-26)$$

The quantity measures the angle between two “vectors” in the subspace $S$ of $L_2(\Phi(x_t) - E[\Phi(x_t)] : t \in T)$. In particular, if two vectors are orthogonal, then $\phi$ is $90^\circ$. If two vectors are in same direction, then $\phi$ is $0^\circ$. Now we are in a position to define the “generalized” correlation coefficient for any two random variables.

**Definition:** Given any two random variables $x$ and $y$, we define a “generalized” correlation coefficient, named *correntropy coefficient* as

$$\eta = \frac{U(x, y)}{\sqrt{U(x, x)U(y, y)}}. \quad (4-27)$$

where $U(x, y)$ is the centered correntropy function for $x$ and $y$, and $U(x, x)$ and $U(y, y)$ are centered correntropy functions for variables themselves.

This definition of correntropy coefficient is in the same spirit of traditional correlation coefficient, where the standard covariance function has been replaced by “generalized”
covariance function, \textit{centered correntropy} Eq. (3–8), and the standard deviation functions
has been replaced by “generalized” deviation function, square root of centered correntropy
with respect to itself Eq. (4–22). However the difference is striking. Conventional
correlation coefficient only measures the second order similarity between two original
random variables, hence it only requires second order statistics in order to attain 0,
i.e., two random variables are uncorrelated. While the correntropy coefficient measures
high order similarity between two original random variables or “second order similarity”
between two nonlinearly transformed random variables in $L_2(\Phi(x_t) - E[\Phi(x_t)] : t \in \mathbb{T})$,
therefore it needs high order statistics to attain 0. In fact, by the property 4 in chapter
2.1, the value of correntropy coefficient is between -1 and 1. If two original random
variables are independent, i.e., two nonlinearly transformed random variables by $\Phi$ are
“uncorrelated” in RKHS induced by kernel function, then the correntropy coefficient
reduces to 0, which means the two vectors in $L_2$ are orthogonal. If two random variables
are the same, the correntropy coefficient is 1, which means the two vectors are in same
direction. This also explains the reason why we use centered correntropy functions,
not correntropy functions, in the definition of correntropy coefficient. The correntropy
coefficient will never attain 0 if correntropy functions are used because $V(x, y)$ is always
greater than 0. The pre-processing of making random processes zero mean is vital to many
conventional statistical signal processing. In our context, since we do not have explicit
knowledge of the mean of nonlinearly transformed random processes, we have to rely on
centered correntropy functions.

Substituting the definition of centered correntropy Eq. (3–8) and approximating the
ensemble average by sample mean, we can obtain an estimate of correntropy coefficient
directly from data,

\[
\hat{\eta} = \frac{1}{N} \sum_{i=1}^{N} \kappa(x_i, y_i) - \frac{1}{N^2} \sum_{i=1}^{N} \sum_{j=1}^{N} \kappa(x_i, y_j) \sqrt{\kappa(0) - \frac{1}{N^2} \sum_{i=1}^{N} \sum_{j=1}^{N} \kappa(x_i, x_j)} \sqrt{\kappa(0) - \frac{1}{N^2} \sum_{i=1}^{N} \sum_{j=1}^{N} \kappa(y_i, y_j)},
\]

(4–28)

where \( \frac{1}{N} \sum_{i=1}^{N} \sum_{j=1}^{N} \kappa(x_i, y_j) \) is called cross information potential between \( x \) and \( y \) [55], \( \frac{1}{N^2} \sum_{i=1}^{N} \sum_{j=1}^{N} \kappa(x_i, x_j) \) and \( \frac{1}{N^2} \sum_{i=1}^{N} \sum_{j=1}^{N} \kappa(y_i, y_j) \) are again the information potential for \( x \) and \( y \) respectively. This observation suggests a connection between the correntropy coefficient and the Cauchy-Schwartz independence measure in Eq. (1–34) because both of them use the information theoretic learning concepts to measure the independence of two random variables. Moreover, the Cauchy-Schwartz independence measure is an approximation to the Kullback-Leibler divergence [55], this also suggests the correntropy coefficient is related to mutual information. These connections need further investigation in the future work.

4.2.2 Representation of RKHS by Centered Correntropy Function

We have analyzed the geometry of \( L_2(\Phi(x_t) - E[\Phi(x_t)] : t \in \mathbb{T}) \), where an inner product of any two vectors is defined in Eq. (4–21) which turns out to be the centered correntropy function for the original random processes. In this section, we proceed to prove that there exist a congruence map between \( L_2(\Phi(x_t) - E[\Phi(x_t)] : t \in \mathbb{T}) \) and RKHS induced by centered correntropy function. Therefore it is sufficient to study the Hilbert space \( L_2(\Phi(x_t) - E[\Phi(x_t)] : t \in \mathbb{T}) \) in this perspective.

First because the centered correntropy function \( U(t, s) \) is non-negative definite, it uniquely determine a reproducing kernel Hilbert space by Moore-Aronszajn Theorem. We can apply Mercer’s theorem to obtain an eigen-decomposition for the centered correntropy function as

\[
U(t, s) = \sum_{k=0}^{\infty} \theta_k \psi_k(t) \psi_k(s) = \langle \Psi(t), \Psi(s) \rangle,
\]

(4–29)
where $\theta_k$ and $\psi_k$ are eigen-values and eigen-functions for the centered correntropy function respectively.

Then we can define a function $g$ on $T$ as the form of

$$g(t) = \sum_{k=0}^{\infty} \theta_k a_k \psi_k(t), \quad (4-30)$$

where the sequence $\{a_k, k = 1, 2, \ldots\}$ satisfies the following condition

$$\sum_{k=0}^{\infty} \theta_k a_k^2 < \infty. \quad (4-31)$$

Let $\mathcal{H}(U)$ be the set composed of functions $g(\cdot)$ which can be represented in the form Eq. (4-30) in terms of eigenfunctions $\psi_k$ and eigenvalues $\theta_k$ of the centered correntropy function $U(t, s)$. Furthermore we might define an inner product of two functions in $\mathcal{H}(U)$ as

$$\langle g_1, g_2 \rangle = \sum_{k=0}^{\infty} \theta_k a_k b_k, \quad (4-32)$$

where $g_1$ and $g_2$ are of form Eq. (4-30) and $a_k, b_k$ satisfy property Eq. (4-31). One might as well show $\mathcal{H}(U)$ is complete. Let $g_n(t) = \sum_{k=0}^{\infty} \theta_k a_k^{(n)} \psi_k(t)$ be a Cauchy sequence in $\mathcal{H}(U)$ such that each sequence $\{a_k^{(n)}, n = 1, 2, \ldots\}$ converges to a limit point $a_k$.

Hence the Cauchy sequence converges to $g(t) = \sum_{k=0}^{\infty} \theta_k a_k \psi_k(t)$ which belongs to $\mathcal{H}(U)$. Therefore $\mathcal{H}(U)$ is a Hilbert space. $\mathcal{H}(U)$ has two important properties which make it a reproducing kernel Hilbert space. First, let $U(t, \cdot)$ be the function on $T$ with value at $s$ in $T$ equal to $U(t, s)$, then by the Mercer’s Theorem eigen-expansion for the covariance function Eq. (4-29), we have

$$U(t, s) = \sum_{k=0}^{\infty} \theta_k a_k \psi_k(s), \quad a_k = \psi_k(t). \quad (4-33)$$

Therefore, $U(t, \cdot) \in \mathcal{H}(U)$ for each $t$ in $T$. Second, for every function $g(\cdot) \in \mathcal{H}(U)$ of form given by Eq. (4-30) and every $t$ in $T$,

$$\langle g, U(t, \cdot) \rangle = \sum_{k=0}^{\infty} \theta_k a_k \psi_k(t) = g(t). \quad (4-34)$$
By the Moore-Aronszajn Theorem, $\mathcal{H}(U)$ is a reproducing kernel Hilbert space with $U(t, s)$ as the reproducing kernel. It follows that

$$\langle U(t, \cdot), U(s, \cdot) \rangle = \sum_{k=0}^{\infty} \theta_k \psi_k(t) \psi_k(s) = U(t, s). \quad (4-35)$$

Thus $\mathcal{H}(U)$ is a representation of the random process $\{x_t : t \in \mathbb{T}\}$ with centered correntropy function $U(t, s)$. One may define a congruence $\mathcal{G}$ form $\mathcal{H}(U)$ onto $L_2(\Phi(x_t) - E[\Phi(x_t)] : t \in \mathbb{T})$ such that

$$\mathcal{G}(U(t, \cdot)) = \Phi(x_t) - E[\Phi(x_t)]. \quad (4-36)$$

The congruence $\mathcal{G}$ can be explicitly represented as

$$\mathcal{G}(g) = \sum_{k=0}^{\infty} a_k \xi_k, \quad (4-37)$$

where the set of $\xi_k$ is an orthogonal random variables belong to $L_2(\Omega, \mathcal{F}, P)$ and $g$ is any element in $\mathcal{H}(U)$ in the form of Eq. (4-30).

This way of proceed from the $L_2(\Phi(x_t) - E[\Phi(x_t)] : t \in \mathbb{T})$ to the RKHS induced by the centered correntropy function is in the same spirit of the method to build RKHS induced by traditional covariance function from the $L_2(x_t, : t \in \mathbb{T})$ [20, 25]. However we now deal with the nonlinearly transformed random processes via the $\Phi$.

### 4.3 Relation Between Two RKHS

There are three spaces we have analyzed so far in this chapter. The first one is the reproducing kernel Hilbert space induced by the data independent kernel function $\kappa(\cdot, \cdot)$, where the nonlinearly transformed random processes reside on the sphere if isotropic kernel functions are used. The inner product in this RKHS is defined as the kernel function $\kappa(\cdot, \cdot)$. The second space is the $L_2(\Phi(x_t) - E[\Phi(x_t)] : t \in \mathbb{T})$ which is built upon the RKHS induced by kernel functions by including all the linear combination of “zero-mean” nonlinearly transformed random process $\{\Phi(x_t) - E[\Phi(x_t)] : t \in \mathbb{T}\}$ and its limit points. An inner product is defined in this space as the form of Eq. (4-21),
which turns out to be the centered correntropy function for the original random processes. Consequently, $L_2 (\Phi(x_t) - E[\Phi(x_t)] : t \in \mathbb{T})$ is complete and thus forms a Hilbert space. The last space is the reproducing kernel Hilbert space induced by the centered correntropy function directly. The inner product is defined as the centered correntropy function, the Mercer kernel. It is proved that there exists an isometric isomorphism (congruence) map between the RKHS induced by the centered correntropy function and the Hilbert space $L_2 (\Phi(x_t) - E[\Phi(x_t)] : t \in \mathbb{T})$. By isometry, it means the inner product definitions for two spaces are the same. By isomorphism, it means there exists a one-to-one mapping between two spaces. It is sufficient to study the RKHS determined by the centered correntropy function by considering the Hilbert space $L_2 (\Phi(x_t) - E[\Phi(x_t)] : t \in \mathbb{T})$. Then the relation among these three spaces become clear. The kernel operator inside the correntropy function induces a reproducing kernel Hilbert space. The expectation operator induces another RKHS which is built upon the first RKHS through the Hilbert space $L_2$ since RKHS by centered correntropy function is congruence to $L_2$.

This interpretation of three spaces involved in centered correntropy functions offers insights into the geometric relationship among the different spaces. Certainly, there are other ways to tackle the task and further investigations are required. First, we applied Mercer's theorem to non-negative definite centered correntropy function to come up with an eigen-decomposition in the previous section. It would further our understanding if we can gain knowledge of the basis. So it is desired that we might also apply what we have propose in section 3.1.2 of explicit construction of RKHS by Gaussian kernel function to build those basis for centered correntropy function explicitly. This will be part of my future work. Second, what we have presented involves high dimensionality, be it in RKHS by kernel function or in RKHS by centered correntropy function. The high dimensionality has its own attractiveness but also poses challenges for data manipulation. So another research direction is to propose a single dimensional nonlinear transformation function such that the correlation function of those nonlinearily transformed functions is
the centered correntropy function, namely to seek a nonlinear function $f$ such that for a given random process \( \{x_t : t \in \mathbb{T}\} \)

$$E[f(x_t) f(x_s)] = U(t, s) = E[\kappa(x_t, x_s)] - E_{x_t} E_{x_s} [\kappa(x_t, x_s)].$$  \hspace{1cm} (4–38)

It is conjectured that this $f$ function will implicitly embed the data distribution. As has been pointed out random processes theorem, any give non-negative definite function determines a Gaussian process such the correlation function for the Gaussian process equals to that non-negative definite function \[74\]. It also suggests that this $f$ function might be related to the gaussianization technique.
One of the most important tasks in statistical analysis is to measure the degree of dependence between random variables. There are numerous measures of dependence in the literature, mostly based on the distance of the joint probability distribution of the data and the product of the marginal probability distributions [80]. Some of the most commonly used measures of bivariate dependence include but are not limited to the correlation coefficient, Kendall’s $\tau$, Spearmans’ $\rho_s$, maximal correlation coefficient, monotone correlation coefficient, and others. The correlation based dependence measures only characterize linear relationship between two random variables. [81] proposed a nonlinear canonical correlation using symmetric nondecreasing functions to quantify nonlinear dependence without any assumption on the distribution of the random variables.

The dependence measure is also closely related to the measure of amount of information that one random variable contained about in the other. Because the more dependent the random variables are, the more information about one ought to be given by the other and vice versa. Several measures based on information theory have been proposed in the literature. For example, the mutual information is one well-known measure which can also be interpreted as the Kullback-Leibler divergence between the joint probability density functions and the product of the marginal densities [82, 83]. It turns out that the Kullback-Leibler divergence or mutual information is a special case of the $\varphi$-divergence measure when a specific convex function $\varphi$ is chosen [84]. On the other hand, [58] generalized Shannon’s mutual information as a special norm in the probability density function space. Silvey’s generalized coefficient [85] uses the Radon-Nikodym derivative of joint distribution of random variables with respect to the product of their marginal distributions. Other dependence measures based on information theory include the relative entropy measure proposed by [86] and others. However Joe’s relative entropy dependence measure, and almost all other entropies fail to be “metric” since they violate
the trianularity rule [87]. In order to overcome this limitation, [87] proposed a metric measure of dependence to specify the degree of dependence in time series data.

Recently there has been considerable work on using functions in reproducing kernel Hilbert space (RKHS) to quantify dependence. [52] introduced kernel dependence functionals in the context of independent component analysis (ICA). The kernelized canonical correlation is a kernelization of canonical correlation analysis with regularization on the functionals. It can also be explained as a particular form of nonlinear canonical correlation where the nonlinear functions have been chosen as functional evaluations in RKHS [88] based on the representer theorem [49]. The kernel generalized variance is an extension of kernelized canonical correlation by estimating the spectral norm of the correlation operator between reproducing kernel Hilbert spaces in the entire spectrum [52]. Instead of using correlation operator in RKHS, [89] proposed the kernel constrained covariance and kernel mutual information based on the covariance operator. The kernel constrained covariance estimates the spectral norm of the covariance operator between RKHSs. It is been proved that the kernel constrained covariance is zero if and only if the original random variables are independent provided that the kernel is universal. The kernel constrained covariance can also be viewed as a maximal correlation coefficient where the function space has been confined to the reproducing kernel Hilbert space. The kernel mutual information incorporates the entire spectrum of the covariance operator between RKHSs and becomes an upper bound of the mutual information estimated with a Parzen window [89]. These dependence measures based on kernel methods have enjoyed much success in independent component analysis [52, 88, 89], quantification of generalized synchronization between chaotic signals [90], and other machine learning application areas.

With the abundance of dependence measures in the literature, one would naturally ask which measure is best in characterizing the dependence information of the data. Rényi proposed a set of postulates which should be fulfilled by a suitable dependence measure [91]. The axiomatic framework by Rényi has drawn much attention ever since.
Many modifications, extensions and enrichments for these postulates have appeared in the literature. One of the major criticisms was that these axioms are too strict for most dependence measures to fulfill. It has been shown by [91] that out of various dependence measures only the maximal correlation coefficient satisfied all these properties. The kernel constrained covariance omits the deterministic dependence and upper bound conditions in the axiomatic framework in order to efficiently estimate the quantity and apply it to independent component analysis [89].

In this chapter, we define a new parametric correntropy function as a novel dependence measure.

5.1 Parametric Correntropy Function

An important observation in the definition of the centered correntropy Eq. (3–8) is that when two random variables are independent \((f_{X,Y}(x, y) = f_X(x)f_Y(y))\) the quantity becomes zero but not vice versa. In order to make it a suitable dependence measure, we modify the definitions of the centered correntropy and correntropy coefficient so that it will satisfy the condition of attaining zero if and only if two random variables are independent.

**Definition (Parametric Centered Correntropy)** Given two random variables \(x\) and \(y\), the parametric centered correntropy is defined as

\[
U_{a,b}(x, y) = E_{x,y}[\kappa(ax, b + y)] - E_x E_y[\kappa(ax, b + y)]
= \iint_{\Omega} \kappa(ax, b + y)(f_{X,Y}(x, y) - f_X(x)f_Y(y))dxdy,
\]

where \(f_{X,Y}(x, y)\) is the joint probability density function (PDF) of random variables \(x\) and \(y\), \(f_X(x)\) and \(f_Y(y)\) are the marginal PDFs respectively, \(\Omega\) is the probability space and \(a\) and \(b\) are parameters in \(\mathbb{R}\).

We present a lemma here stating that the parametric centered correntropy is zero for all \(a\) and \(b\) if and only if \(x\) and \(y\) are independent. First, we employ the Fourier transform
for the shift-invariant real valued kernel function as

$$\kappa(z) = \int e^{i\alpha z} \mu(d\alpha), \quad (5-1)$$

where $\mu$ is a positive bounded measure. For simplicity we will also assume that the probability measure $\mu$ has moments of all orders so that $\kappa$ is infinitely differentiable. If the tail of $\mu$ decays sufficiently fast $\kappa$ can be extended to the entire line.

**Lemma (Independence):** Suppose the measure $\mu$ defining the kernel $\kappa$ has support in the entire line. Then $U_{a,b}(x, y) = 0$ for all $a$ and $b$ if and only if $x$ and $y$ are independent.

**Proof:** The sufficient condition is straightforward. Assume $U_{a,b}(x, y) = 0$ for all $a$ and $b$, re-write the definition of the parametric centered correntropy as

$$U_{a,b}(x, y) = \int \int \Omega \kappa(ax - y - b) \{dF_{X,Y}(x, y) - dF_X(x)dF_Y(y)\}$$

$$= \int \int \Omega \kappa(ax - y - b)dQ(x, y), \quad (5-2)$$

where $F_{X,Y}(x, y)$ is the joint cumulative distribution function (CDF), and $F_X(x)$ and $F_Y(y)$ are the marginal CDFs for $x$ and $y$ respectively. Need to show that $dQ(x, y) = 0$.

$U_{a,b}(x, y) = 0$ for all $a$ and $b$ means $\int \int e^{i\alpha(x - y)}dQ(x, y)e^{-i\alpha b}\mu(d\alpha) = 0$ for all $a$ and $b$, in particular for all $b$. Hence $\int \int e^{i\alpha(x - y)}dQ(x, y) = 0$ $\mu$ almost all $\alpha$. Since support $\mu = \mathbb{R}$, this holds for all $\alpha$ and $a$. This is easily written as $\int e^{i\alpha x + \beta y}dQ(x, y) = 0$ for all $\alpha$ and $\beta$. Conclusion $dQ = 0$. ■

We can also parameterize the correntropy coefficient.
Definition (Parametric Correntropy Coefficient) Given two random variables \(x\) and \(y\), the parametric correntropy coefficient is defined as

\[
\eta_{a,b}(x, y) = \frac{E_{x,y}[\kappa(ax, b + y)] - E_x E_y[\kappa(ax, b + y)]}{\sqrt{E[\kappa(ax, ax)] - E_x E_x[\kappa(ax, ay)] E[\kappa(b + y, b + y)] - E_y E_y[\kappa(b + x, b + y)]}} \\
= \frac{\int_{\Omega} \kappa(ax, b + y)(f_{X,Y}(x, y) - f_X(x) f_Y(y)) dx dy}{\sqrt{\kappa(0) - \int_{\Omega} \kappa(ax, ay) f_X(x) f_Y(y) dx dy}},
\]

where \(\kappa(0)\) is the value of kernel function when its argument is zero. The absolute value of the parametric correntropy coefficient is bounded by 1.

Proof (Boundedness of parametric correntropy coefficient): By the eigen-decomposition of the kernel function Eq. (4–1), we have

\[
|E_{x,y}[\kappa(ax, b + y)] - E_x E_y[\kappa(ax, b + y)]| \\
= \left| E_{x,y} \left[ \sum_{k=0}^{\infty} \lambda_k \varphi_k(ax) \varphi_k(b + y) \right] - E_x E_y \left[ \sum_{k=0}^{\infty} \lambda_k \varphi_k(ax) \varphi_k(b + y) \right] \right| \\
= \sum_{k=0}^{\infty} \lambda_k \left| E[\varphi_k(ax) \varphi_k(b + y)] - E[\varphi_k(ax)]E[\varphi_k(b + y)] \right| \\
= \sum_{k=0}^{\infty} \lambda_k \left| E\{\varphi_k(ax) - E[\varphi_k(ax)]\}(\varphi_k(b + y) - E[\varphi_k(b + y)]) \right| \\
\leq \sum_{k=0}^{\infty} \lambda_k \sqrt{\text{var}(\varphi_k(ax))\text{var}(\varphi_k(b + y))} \left( |\text{cov}(XY)| \leq \sqrt{\text{var}(X)\text{var}(Y)} \right) \\
\leq \left( \sum_{k=0}^{\infty} \lambda_k \text{var}(\varphi_k(ax)) \right)^{1/2} \left( \sum_{k=0}^{\infty} \lambda_k \text{var}(\varphi_k(b + y)) \right)^{1/2} \quad (\text{Cauchy – Schwartz inequality})
\]

(5–4)

where

\[
\sum_{k=0}^{\infty} \lambda_k \text{var}(\varphi_k(ax)) = \sum_{k=0}^{\infty} \lambda_k \left( E[\varphi_k^2(ax)] - E^2[\varphi_k(ax)] \right).
\]

(5–5)
The first term of Eq. (5–5) is
\[ \sum_{k=0}^{\infty} \lambda_k E[\varphi_k^2(ax)] = E[\sum_{k=0}^{\infty} \lambda_k \varphi_k^2(ax)] = \kappa(0), \]
and the second term of Eq. (5–5) is
\[ \sum_{k=0}^{\infty} \lambda_k E^2[\varphi_k(ax)] = \sum_{k=0}^{\infty} \lambda_k \left( \int_{\Omega} \varphi_k(ax)f_X(x)dx \right)^2 \\
= \sum_{k=0}^{\infty} \lambda_k \left( \int_{\Omega} \varphi_k(ax)f_X(x)dx \right) \left( \int_{\Omega} \varphi_k(ay)f_X(y)dy \right) \\
= \int_{\Omega} \left( \sum_{k=0}^{\infty} \lambda_k \varphi_k(ax) \varphi_k(ay) \right) f_X(x)f_X(y)dx dy \\
= \int_{\Omega} \kappa(ax, ay) f_X(x)f_X(y) dx dy \\
= E_x E_y [\kappa(ax, ay)]. \]

Therefore,
\[ \sum_{k=0}^{\infty} \lambda_k \text{var}(\varphi_k(ax)) = \kappa(0) - \int_{\Omega} \kappa(ax, ay)f_X(x)f_X(y)dx dy. \tag{5–6} \]

Likewise
\[ \sum_{k=0}^{\infty} \lambda_k \text{var}(\varphi_k(b + y)) = \kappa(0) - \int_{\Omega} \kappa(b + x, b + y)f_Y(x)f_Y(y)dx dy. \tag{5–7} \]

Combining Eq. (5–4), Eq. (5–6) and Eq. (5–7) together, we obtain
\[ |E_x,y[\kappa(ax, b + y)] - E_x E_y[\kappa(ax, b + y)]| \leq \left( \kappa(0) - \int_{\Omega} \kappa(ax, ay)f_X(x)f_X(y)dx dy \right)^{1/2} \left( \kappa(0) - \int_{\Omega} \kappa(b + x, b + y)f_Y(x)f_Y(y)dx dy \right)^{1/2}. \]

Hence the absolute value of the parametric correntropy coefficient is bounded by 1. □

5.2 Correntropy Dependence Measure

Based on the parametric correntropy coefficient developed in the previous section, we formulate a novel dependence measure for two random variables. Rényi gave a set
of fundamental principles that a measure of statistical dependence $Q(x, y)$ between two random variables $x$ and $y$ should satisfy. These include

1. $Q(x, y)$ is well defined,
2. $0 \leq Q(x, y) \leq 1$,
3. $Q(x, y) = 0$ if and only if $x$ and $y$ are independent,
4. $Q(x, y) = 1$ if and only if $y = f(x)$ or $x = g(y)$, where $f$ and $g$ are Borel measurable functions.

Rényi showed that one measure satisfying these constraints is

$$Q(x, y) = \sup_{f,g} \text{corr}(f(x), g(y)), \quad (5–8)$$

where $f(x)$ and $g(y)$ must have finite positive variance, and $f, g$ are Borel measurable.

We proposed a dependence measure based on the parametric correntropy coefficient which satisfies all the desirable properties above.

**Definition (Correntropy Dependence Measure)** Given two random variables $x$ and $y$, the Correntropy Dependence Measure is defined as

$$\Gamma(x, y) = \sup_{a,b} |\eta_{a,b}(x, y)|, \quad (5–9)$$

where $\eta_{a,b}(x, y)$ is the parametric correntropy coefficient Eq. (5–3).

The correntropy dependence measure is a suitable statistical measure which fulfills all the fundamental conditions listed by Rényi.

**Proof:** First the measure Eq. (5–9) is well defined. Since $|\eta_{a,b}(x, y)|$ is between 0 and 1 as proved above, certainly the supreme of $|\eta_{a,b}(x, y)|$ is also bounded by 0 and 1. The independence condition is also trivial. If $x$ and $y$ are independent, $|\eta_{a,b}(x, y)| = 0$, therefore $\sup_{a,b} |\eta_{a,b}(x, y)| = 0$. Since

$$0 \leq |\eta_{a,b}(x, y)| \leq \sup_{a,b} |\eta_{a,b}(x, y)|,$$
if \( \sup_{a,b} |\eta_{a,b}(x,y)| = 0 \), then \( |\eta_{a,b}(x,y)| = 0 \). Therefore \( x,y \) are independent. To check the condition 4, consider the derivation in the boundedness proof Eq. (5–4). The equalities achieve if and only if when

\[
\varphi_k(ax) - E[\varphi_k(ax)] = \alpha(\varphi_k(b + y) - E[\varphi_k(b + y)]), \quad \forall k, \tag{5–10}
\]

and,

\[
E[(\varphi_k(ax) - E[\varphi_k(ax)])^2] = \beta E[(\varphi_k(b + y) - E[\varphi_k(b + y)])^2], \quad \forall k. \tag{5–11}
\]

In fact, both conditions Eq. (5–10) and Eq. (5–11) are equivalent. From the condition Eq. (5–10), we obtain

\[
\varphi_k(ax) = B_k\varphi_k(b + y) + A_k, \quad \forall k \tag{5–12}
\]

where \( B_k \) and \( A_k \) are two constant. Let

\[
f(x) = \sum_{k=0}^{\infty} C_k\varphi_k(ax) = \sum_{k=0}^{\infty} C_k(B_k\varphi_k(b + y) + A_k) = g(y), \tag{5–13}
\]

then \( x = f^{-1}(g(y)) \) since functional \( \varphi_k \) is continuous and invertible being the eigen-function of the positive definite kernel. Therefore we conclude the proof. ■

Unlike the statistical measure proposed by Rényi Eq. (5–8), the correntropy dependence measure searches the parameter space for a supreme instead of the entire function space. Compared to the kernel constrained covariance presented in [89], our measure does not employ regularization technique because our optimization space is the one dimensional parameter space. However the kernel constrained covariance operates on the high-dimensional (possibly infinite dimensional) functional space, therefore regularization is mandatory in order to make solution achievable. By selecting different values of parameters \( a \) and \( b \), the correntropy dependence measure offers a different scale value of dependence between random variables \( x \) and \( y \). The idea is similar to the concept of wavelet which provides multi-resolution information about signal frequency.
Principal component analysis (also known as the Karhunen-Loève transformation in communication theory) is a powerful tool for feature extraction and data dimensionality reduction in statistical pattern recognition and signal processing. It can be easily performed by eigen-decomposition of the standard covariance matrix or by adaptive algorithms that estimate principal components [92]. Principal component analysis or PCA is really an affine transformation of the coordinate system such that the rate of decrease of data variance is maximized. The projections of the data onto the new coordinate system are called principal components. These projections represent the data optimally in a least-square sense. In feature extraction, PCA transforms the data in such a way that a small number of principal components can represent the data while retaining most of the intrinsic variance of the data. These are sometimes called factors or latent variables of the data [93].

While PCA yields a smaller dimensional linear subspace that best represents the full data according to a minimum-square-error criterion, it might be a poor representation if the data structure is non-Gaussian. Hence nonlinear component analysis may be needed. There have been numerous attempts to define nonlinear components analysis in the latest decades. Nonlinear PCA is generally seen as a nonlinear generalization of standard PCA [92, 93]. The principal component is generalized from straight lines to curves. Principal curves were proposed by Hastie [94] to define local directions that pass through the high density parts of the data set. The principal curves are found through an iterative algorithm that minimizes the conditional expectation of projections on the curves. Kramer presented a nonlinear PCA based on auto-associative neural networks. The auto-associative network performs identity mapping from the input data to the output by minimizing the square error [95]. Recently, Schölkopf et al applied kernel methodology to obtain a nonlinear form of PCA [50]. This so called Kernel PCA is one of the kernel-based
learning algorithms reviewed in Chapter 1.3. The Kernel PCA nonlinearly maps the
original input data into an infinite dimensional reproducing kernel Hilbert space by the
data independent kernel and solves the eigen-decomposition of the Gram matrix of the
input data in a high-dimensional feature space. The Gram matrix has a dimension given
by the number of samples $N$. The data projections onto the principal directions of the
Gram matrix, i.e. the inner product in feature space, are carried out by means of kernel
functions in the input space. While the utilization of Mercer kernels provides a tractable
way to compute principal components in the high-dimensional feature space, there are
still problems of interpretation and computation of the large dimensional Gram matrix.
Indeed, the number of eigenfunctions of the Gram matrix is dependent on the number of
data samples $N$, not the size of the data space $L$. Moreover computing Gram matrices for
millions of samples in a small, let us say, two dimensional space becomes wasteful.

In this chapter, we propose a new nonlinear PCA technique based on the correntropy
function, called correntropy PCA. The correntropy function quantifies the similarity
between the $L$ different components of the $L$ dimensional input data vector (or the
time structure in a time series) using the statistical data distribution. The correntropy
also utilizes a kernel methodology, but in a different form: by applying the kernel to
pairs of data vector components, a random vector (or stochastic process) is nonlinearly
transformed into a high dimensional function space where the similarity between the
components of the transformed random variables (or stochastic process) can be measured
by the conventional covariance function. The eigen-decomposition of the covariance
of the transformed data yields the principal directions of the nonlinearly transformed
data. These linear principal directions in feature space correspond to nonlinear principal
directions in the input space. These projections can be efficiently computed by utilizing
the correntropy function. That means, if one has one million samples in a two dimensional
space, it is only necessary to solve a two dimensional eigenvector problem on a matrix
whose entries are computed from one million samples. In many applications this is a
tremendous computational saving.

Given a set of zero mean vector observations $x_j$, $j = 1, \ldots, N$, $x_j \in \mathbb{R}^L$, $\sum_{j=1}^{N} x_j = 0$, correntropy PCA seeks a direction in the feature space such that the variance of the data
projected onto this direction is maximized. Unlike the kernel method which transforms
data into a feature space sample by sample, correntropy PCA maps data component-wise
into a feature space, i.e., the RKHS associated with the centered correntropy function. By
the equation Eq. (4–29),

$$
\Psi : \mathbb{R}^L \mapsto F
$$

$$
x \mapsto [\Psi(x_1), \Psi(x_2), \cdots, \Psi(x_L)],
$$

where $x_i$ denotes the $i^{th}$ component of the original input data sample $x$. This nonlinear
mapping transforms the component-wise data into a high dimensional RKHS which
is associated with the centered correntropy function. By the definition of centered
correntropy function, we have

$$
<\Psi(x_i), \Psi(x_j)> = U(x_i, x_j) = E_{x_i,x_j}\{\kappa(x_i, x_j)\} - E_{x_i}E_{x_j}\{\kappa(x_i, x_j)\}
$$

$$
= \frac{1}{N} \sum_{k=1}^{N} \kappa(x_{ik}, x_{jk}) - \frac{1}{N^2} \sum_{k=1}^{N} \sum_{m=1}^{N} \kappa(x_{ik}, x_{jm}), \forall \ i, j = 1, 2, \ldots, L,
$$

where $x_{ik}$ is the $i^{th}$ component of the $k^{th}$ input data sample. The expectation runs over all
the data samples.

Then the covariance matrix of the transformed data in the feature space is given by

$$
C = \frac{1}{L} \sum_{i=1}^{L} \Psi(x_i)\Psi^T(x_i)
$$

We now have to find the eigenvalues $\lambda \geq 0$ and non-zero eigenvectors satisfying

$$
Cq = \lambda q.
$$
All the solutions $q$ must lie in the span of $\Psi(x_1), \ldots, \Psi(x_L)$, i.e., we $q$ is the form of linear combination of all the $\Psi(x_1), \ldots, \Psi(x_L)$,

$$q = \sum_{i=1}^{L} \beta_i \Psi(x_i) \quad (6-2)$$

And we may instead consider the set of equations,

$$< \Psi(x_k), Cq > = < \Psi(x_k), \lambda q >, \forall k = 1, \ldots, L. \quad (6-3)$$

Combining equations Eq. (6–2) and Eq. (6–3), we get

$$< \Psi(x_k), \frac{1}{L} \sum_{j=1}^{L} \Psi(j) \Psi^T(j) \cdot \sum_{i=1}^{L} \beta_i \Psi(x_i) >$$

$$= \frac{1}{L} \sum_{j=1}^{L} \sum_{i=1}^{L} \beta_i < \Psi(x_k), \Psi(x_j) > < \Psi(x_j), \Psi(x_i) >$$

$$= \lambda \sum_{i=1}^{L} \beta_i < \Psi(x_k), \Psi(x_i) >, \forall k = 1, \ldots, L. \quad (6-4)$$

By equation Eq. (6–1), we can define an $L \times L$ centered correntropy matrix $U$ by

$$U_{ij} : = E[\kappa(x_i, x_j)] - E_{x_i} E_{x_j}\kappa(x_i, x_j) \quad (6-5)$$

$$= \frac{1}{N} \sum_{k=1}^{N} \kappa(x_{ik}, x_{jk}) - \frac{1}{N^2} \sum_{k=1}^{N} \sum_{m=1}^{N} \kappa(x_{ik}, x_{jm}),$$

$$\forall \ i, j = 1, 2, \ldots, L.$$

Let $k$ in Eq. (6–4) runs from 1 to $L$, and write the result in matrix form, we can get

$$U^2 \beta = L \lambda U \beta, \quad (6-6)$$

where $\beta$ denotes the column vector with entries $\beta_1, \ldots, \beta_L$. It can be shown that the solutions of equation Eq. (6–6) are equivalent to the solutions to the following eigenvalue problem,

$$U \beta = L \lambda \beta, \quad (6-7)$$

for nonzero eigenvalues.
For the purpose of principal component extraction, we need to compute the projections onto the eigenvectors $\mathbf{q}$ in the feature space. Let $\mathbf{x}$ be a test point, the projection of $\mathbf{x}$ onto the principal direction mapped back to input space is given by

$$P(\mathbf{x}) = \sum_{i=1}^{L} \beta_i \frac{1}{N} \sum_{j=1}^{N} \kappa(\mathbf{x}_{ij}, \mathbf{x}_i), \quad (6-8)$$

this is so called a nonlinear principal component.

In summary, we need to take the following steps to compute the nonlinear principal components: (1) compute the correntropy matrix $V$ by equation Eq. (6-5), where the expected value is substituted by the average, (2) compute its eigenvectors and eigenvalues through SVD, and (3) compute the projections of a test point onto the eigenvectors by Eq. (6–8).

We will present two experimental results to show the effectiveness of correntropy PCA in finding nonlinear principal directions. The first experiment compares the standard linear PCA and correntropy PCA to extract features from a two dimensional mixture of Gaussian distributed data. Specifically, the probability density function is a mixture of Gaussian modes with the following form

$$f(\mathbf{x}) = 1/2(\mathcal{N}(\mathbf{m}_1, \Sigma_1) + \mathcal{N}(\mathbf{m}_2, \Sigma_2)), $$

where $\mathcal{N}(\mathbf{m}_1, \Sigma_1)$ and $\mathcal{N}(\mathbf{m}_2, \Sigma_2)$ are two Gaussian distributed data with the mean vectors and variance matrices given by

$$\begin{align*}
\mathbf{m}_1 &= \begin{bmatrix} -1 \\ -1 \end{bmatrix}, & \Sigma_1 &= \begin{bmatrix} 1 & 0 \\ 0 & 0.1 \end{bmatrix}, \\
\mathbf{m}_2 &= \begin{bmatrix} 1 \\ 1 \end{bmatrix}, & \Sigma_2 &= \begin{bmatrix} 0.1 & 0 \\ 0 & 1 \end{bmatrix}.
\end{align*}$$

In Fig. 6-1, we plot the contours of the data and of the largest eigenvalue directions produced by linear PCA and correntropy PCA respectively. 200 samples are used and
Figure 6-1. Linear PCA versus correntropy PCA for a two-dimensional mixture of Gaussian distributed data

kernel size is chosen to be 2. The result confirms that linear PCA only provides the linear directions that maximizes the variance. But since the underlying data is a mixture of two Gaussian modes, linear PCA fails to consider the directions of the individual modes but only averages these directions. On the contrary, correntropy PCA is more tuned to the underlying structure of the data in the input space. Correntropy PCA generates a nonlinear principal direction that follows locally the directions of the individual modes so that the variance of principal component projected onto this nonlinear curve is maximized. The experiment shows that correntropy PCA is superior in describing the underlying structure of the data when compared to the linear PCA method.

Our second experiment compared the kernel PCA, proposed by Schölkopf et al in [50], with correntropy PCA. We use the same experiment setup as in [50] in order to illustrate the performance of correntropy PCA. The data is two-dimensional with three clusters (Gaussian distribution with standard deviation 0.1). The number of data sample and the kernel size are chosen to be 90 and 0.1 respectively. Since the number of principal
components for kernel PCA depends on the number of data samples, there are many eigen-directions in feature space that are difficult to identify in the input space, so we plot the two principal components with the largest eigenvalues from kernel PCA. However the number of principal components for correntropy PCA is equal to the dimension of input space, so there is no ambiguity. Fig. 6-2 shows that both kernel PCA and correntropy PCA can extract the nonlinear principal components form the data. While kernel PCA tends to find the local structure for a given data set as the contours circle around different data clusters suggest, correntropy PCA seeks the underlying global structure of the data set. The contour in the left bottom plot shows that correntropy PCA can be tuned to the data structure by changing the kernel size in the Gaussian kernel, and locate the principal direction.

In experiments comparing the performance of correntropy PCA with standard linear PCA and kernel PCA for nonlinear feature extraction, we found two advantages of our method. First, correntropy PCA can be more tuned to the underlying data structure
than linear PCA so that it can extract the nonlinear principal components from the data, very much like principal curves. There is no ambiguity since the number of nonlinear principal components is the same as the dimensionality of the input space. In kernel PCA it is very difficult to choose the eigen-directions if we can not visualize the data, since the eigenvectors project locally to the input space. Therefore, it is not easy to separate major and minor components. Second, correntropy PCA has a tremendous computational complexity advantage over kernel PCA. For example, in the second simulation, we only need to compute an eigen-decomposition for a $2 \times 2$ matrix using correntropy PCA while we have to do eigen-decomposition for a $90 \times 90$ matrix using kernel PCA. As the training set increases, the computational complexity of kernel PCA will increase dramatically but the size of the correntropy matrix remains the same.

In this chapter we applied correntropy concepts in principal component analysis. The approach is based on finding the eigenvectors of the centered correntropy matrix (same as the dimension of the data) unlike the Gram matrix used by other kernel methods, where the dimension of the Gram matrix is dependent on the number of the data. Yet, the final principle curves we get using this method adequately covers the data in the direction of maximum spread (variance in the feature space). Since we are dealing with a finite dimensional matrix, we get a number of principle curves equal to the dimension of the data space, and at the same time the computational complexity is drastically reduced compared to the kernel methods. In general this approach offers a new method of analyzing data. The study also suggests that the concept of correntropy can be used for de-correlating the data in the feature space (whitening), which can be applied in the context of independent component analysis. The future research will apply correntropy PCA to real data problem and also compare with other nonlinear principal component analysis methods.
7.1 Introduction

Pitch, or the fundamental frequency $F_0$, is an important parameter of speech signals. Accurate determination of pitch plays a vital role in acoustical signal processing and has a wide range of applications in related areas such as coding, synthesis, speech recognition and others. Numerous pitch determination algorithms (PDA) have been proposed in the literature $[96]$. In general, they can be categorized into three classes: time-domain, frequency-domain, and time-frequency domain algorithms.

Time-domain PDAs operate directly on the signal temporal structure. These include but are not limited to zero-crossing rate, peak and valley positions, and autocorrelation. The autocorrelation model appears to be one of the most popular PDAs for its simplicity, explanatory power, and physiological plausibility. For a given signal $x_n$ with $N$ samples, the autocorrelation function $R(\tau)$ is defined as

$$R(\tau) = \frac{1}{N} \sum_{n=0}^{N-1} x_n x_{n+\tau}, \quad (7-1)$$

where $\tau$ is the delay parameter. For dynamical signals with changing periodicities, a short-time window can be included to compute the periodicities of the signal within the window ending at time $t$ as

$$R(\tau, t) = \frac{1}{N} \sum_{n=0}^{N-1} x_n x_{n+\tau} w_t, \quad (7-2)$$

where $w_t$ is an arbitrary causal window that confines the autocorrelation function into a neighborhood of the current time. Other similar models can be obtained by replacing the multiplication by subtraction (or excitatory by inhibitory neural interaction) in the autocorrelation function such as the average magnitude difference function (AMDF) $[97]$. Cheveigné proposed the squared difference function (SDF) in $[98]$ as

$$\text{SDF}(\tau) = \frac{1}{N} \sum_{n=0}^{N-1} (x_n - x_{n+\tau})^2. \quad (7-2)$$
The weighted autocorrelation uses an autocorrelation function weighted by the inverse of an AMDF to extract pitch from noisy speech [99]. All these PDAs based on the autocorrelation function suffer from at least one unsatisfactory fact: the peak corresponding to the period for a pure tone is rather wide [100, 101]. This imposes greater challenge for multiple $F0$ estimation since mutual overlap between voices weakens their pitch cues, and cues further compete with cues of other voices. The low resolution in pitch estimation results from the fundamental time-frequency uncertainty principle [102]. To overcome this drawback, Brown et al. presented a “narrowed” autocorrelation function to improve the resolution of the autocorrelation function for musical pitch extraction [103]. The “narrowed” autocorrelation function includes terms corresponding to delays at $2\tau$, $3\tau$, etc., in addition to the usual term with delay $\tau$ as

$$S_L(\tau) = \frac{1}{N} \sum_{n=0}^{N-1} (x_n + x_{n+\tau} + x_{n+2\tau} + \cdots + x_{n+L\tau})^2. \quad (7-3)$$

However it requires an increase in the length of the signal and less precision in time. It also requires the $a$ priori selection of the number of delay terms $L$.

Frequency-domain PDAs estimate pitch by using the harmonic structure in the short-time spectrum. Frequency-domain methodologies include component frequency ratios, filter-based methods, cepstrum analysis and multi-resolution methods. Pitch determination algorithms such as harmonic sieve [104], harmonic product spectrum [105], sub-harmonic summation [106], and subharmonic to harmonic ratio [107] fall into this category. Most frequency-domain pitch determination methods apply pattern matching [108]. Others use nonlinear or filtering preprocessing to generate or improve interpartial spacing and fundamental component cues. The frequency-domain PDAs have the advantage of efficient implementation with fast Fourier transform and theoretical strength of Fourier analysis. However one weakness is that they rely on the shape and size of the analysis window. Selection and adjustment of analysis window remain a problem in estimation.
The time-frequency approach splits the signal over a filter-bank, applies time-domain methods to each channel waveform, and the results are aggregated over channels. The summary, or “pooled”, autocorrelation functions across all channels provides pitch information of the signal. Licklider first presented this idea as a pitch perception model [109]. Later Lyon and Slaney further developed the methodology and called it correlogram [110, 111]. The correlogram is the first stage processor in a computational auditory scene analysis (CASA) system [112]. It has also been incorporated into a neural oscillator to segregate double vowels and multipitch tracking [101, 113]. The strength of correlogram in pitch estimation is that different frequency channels corresponding to different signal sources of different pitches can be separated, which makes it useful in multipitch estimation [101, 114]. Also individual channel weighting can be adapted to compensate for amplitude mismatches between spectral regions [115].

On the other hand, autocorrelation and power spectrum based pitch determination algorithms mentioned above only characterizes second-order statistics. In many applications where non-Gaussianities and nonlinearities are present, these second-order statistical methodologies might fail to provide all the information about the signals under study. Higher-order statistics have been used in pitch determination. Moreno et al. applied higher-order statistics to extract pitch from noisy speech [116]. But only diagonal third order cumulants were used for simplicity and computational efficiency which is given by

\[ c(k) = \frac{1}{N} \sum_{n=0}^{N-1} x_n x_{n+k}, \quad k = 0, \ldots, N - 1. \]

And pitch is found by applying autocorrelation function to the cumulants \( c(k) \),

\[ G(\tau) = \frac{1}{2N} \sum_{k=-(N-1)}^{N-1} c(k)c(k + \tau). \] (7-4)
In this chapter, we propose a new pitch determination algorithm based on the correntropy function defined in the previous chapters,

\[
V(\tau) = \frac{1}{N} \sum_{n=0}^{N-1} \kappa(x_n, x_{n+\tau}).
\]  

(7-5)

The proposed pitch determination method is applied after the acoustical signal is processed by an equivalent rectangular bandwidth (ERB) filter bank in the time domain. The ERB filter bank acts as a cochlear model to transform a one dimensional acoustical signal into a two dimensional map of neural firing rate as a function of time and place \[111\]. The correntropy function for each channel is calculated and the summation across all the channels provides the pitch information. As a novel “self-similarity” measure, correntropy is able to offer much better resolution than the conventional autocorrelation function in pitch estimation. Moreover, our pitch determination algorithm can segregate double vowels without applying any complex model such as a neural oscillator \[101\].

7.2 Pitch Determination based on Correntropy

The structure of the correntropy definition seems very appropriate to quantify similarity in time, in a manner that is biologically plausible, that is, it can be implemented at neuronal level. Its argument is sensitive to differences in time instances as correlation, but instead of being linear across differences, there is an intermediate nonlinear transformation that gives more emphasis to values that are closer together. Neurons are known to be very sensitive to time differences, but their highly nonlinear response may also emphasize similarity at the close range. In the context of pitch determination, the correntropy function might as well estimate the pitch information of the signal similar to the autocorrelation function. However, compared to the autocorrelation function model, our pitch determination algorithm based on the correntropy function offers much better resolution and enhances the capacity of estimating multiple pitches. Since correntropy creates many different harmonics of each resonance present in the original time series due to the nonlinearity of the kernel function, it may also be useful for perceptual pitch
determination. To illustrate this, we compare the conventional autocorrelation Eq. (7–1), narrowed correlation Eq. (7–3) and correntropy functions Eq. (7–5) for a simple sinusoid signal. Fig. 7-1 plots three functions in the lag with respect to the delay lag in the time domain. All three functions are able to peak at the same delay lag corresponding to the period of the sinusoid signal. However it is evident that the peaks obtained from the correntropy function are much narrower and sharper than those from the conventional autocorrelation and narrowed autocorrelation functions. In Fig. 7-2, we present the Fourier transform of each function. The ordinary autocorrelation function only exhibits one harmonic and the narrowed autocorrelation produces 10 harmonics which is equal to the number of terms $L$ used in Eq. (7–3). The correntropy function places even more energy at higher harmonics in frequency. The narrowness of correntropy function in the time domain implies the rich harmonics present in the frequency domain. This is due to the nonlinear exponential function included in the definition of the correntropy function.

It should also be noticed that there is a connection between the correntropy function Eq. (7–5) and the square difference function Eq. (7–2). The correntropy function also uses inhibitory neural interaction model instead of excitatory one with a Gaussian kernel function, but it nonlinearly transforms the subtraction of the signals by the exponential
Figure 7-2. Fourier transform of autocorrelation, narrowed autocorrelation with \( L = 10 \) and correntropy functions of a sinusoid signal.

function. From another perspective, the correntropy function includes the scaled square different function as an individual term for \( k = 1 \) in the summation of Eq. (3–11). But it contains more information with other higher-order moment terms.

Our pitch determination algorithm first uses cochlear filtering to peripherally process the speech signal. This is achieved by a bank of 64 gammatone filters which are distributed in frequency according to their bandwidths [117]. The impulse response of a gammatone filter is defined as

\[
q(t) = t^{n-1}e^{-2\pi at} \cos(2\pi f_0 t + \psi),
\]

where \( n \) is the filter order with center frequency at \( f_0 \) Hz, \( \psi \) is phase, and \( a \) is bandwidth parameter. The bandwidth increases quasi-logarithmically with respect to the center frequency. The center frequencies of each filter are equally spaced on the equivalent rectangular bandwidth scale between 80 Hz and 4000 Hz [118]. This creates a cochleagram, which is a function of time lag along the horizontal axis and cochlear place, or frequency, along the vertical axis. The cochlear separates a sound into broad frequency channels while still contains the time structure of the original sound. It has served as a peripheral
Figure 7-3. Correlogram (top) and summary (bottom) for the vowel /a/.

The periodicity analysis is done by computing the correntropy function at the output of each cochlear frequency channel,

\[
V_i(\tau) = \frac{1}{N} \sum_{n=0}^{N-1} \kappa(x^i_n, x^i_{n+\tau}),
\]

(7-6)

where \(i\) stands for channel number and \(z_n\) is the cochlear output. The kernel bandwidth is determined using Silverman’s rule [120]. The time lag \(\tau\) is chosen long enough to include the lowest expected pitch. Generally it is set at least 10ms throughout the paper. In this way, a picture is formed with horizontal axis as correntropy lags and vertical axis as cochlear frequency. We name it correntropy-gram, which literally means “pictures of correntropy”. If a signal is periodic, strong vertical lines at certain correntropy lags appear in the correntropy-gram indicating times when a large number of cochlear
channels are firing synchronously. While the horizontal bands signify different amounts of energy across frequency regions. The correntropy-gram is similar to the correlogram in structure but different in content. In order to reduce the dynamic range for display in the correntropy-gram, the correntropy functions should be normalized such that the zero lag value is one as given by the following formula,

\[
C_i(\tau) = \frac{\frac{1}{N} \sum_{n=0}^{N-1} \kappa(x_{n}^i, x_{n+\tau}^i) - \frac{1}{N^2} \sum_{n,m=0}^{N-1} \kappa(x_{n}^i, x_{m+\tau}^i)}{\sqrt{V(0) - \frac{1}{N^2} \sum_{n,m=0}^{N-1} \kappa(x_{n}^i, x_{m}^i)}} \cdot \sqrt{V(0) - \frac{1}{N^2} \sum_{n,m=0}^{N-1} \kappa(x_{n+\tau}^i, x_{m+\tau}^i)}},
\]

where \(V(0)\) is the value of correntropy when lag \(\tau = 0\). The numerator is called the centered correntropy which takes out the mean value of the transformed signal in the RKHS. \(C(\tau)\) is also called correntropy coefficient that has been applied to detect nonlinear dependence among multichannel biomedical signals [121].

Figure 7-4. Autocorrelation (top) and summary (bottom) of third order cumulants for the vowel /a/. 
Figure 7-5. Narrowed autocorrelation (top) and summary (bottom) for the vowel /a/.

In order to emphasize pitch related structure in the correntropy-gram, the correntropy functions are summed up across all the channels to form a “pooled” or “summary” correntropy-gram,

$$W(\tau) = \sum_i V_i(\tau).$$

The summary correntropy-gram measures how likely a pitch would be perceived at a certain time lag. The pitch frequency can be obtained by inverting the time delay lag. In our experiment, the summary of correntropy functions is first normalized by subtracting the mean and dividing by the maximum absolute value. The position of pitch can be picked by various peak-picking algorithms to identify local maximum above the pre-defined threshold. Here we calculate the first derivative and mark the position when the value changes from positive to negative as a local maximum.

Compared to the conventional correlogram model [101, 119], [122], our pitch detector is able to locate the same period information as the correlogram, but has much narrower
peaks. Hence the proposed method enhances the resolution of pitch determination. Furthermore, since the correlogram estimates the likelihood that a pitch exists at a certain time delay, the summary correlogram may generate other “erroneous” peaks besides the one corresponding to the pitch period [111]. while the summary correntropy-gram suppresses values that are dissimilar at all other time delays by the exponential decay of the Gaussian function and only peaks at the one corresponding to the pitch period. For mixtures of concurrent sound sources with different fundamental frequencies, the summary correlogram usually fails to detect multiple pitches without further nonlinear post-processing. But the summary correntropy-gram is able to show peaks at different periods of each source. These characteristics of the proposed method suggest a superiority of the correntropy function over the autocorrelation function in pitch determination.

Moreover, the computational complexity of our method, whether the correntropy function Eq. (7–6) or the correntropy coefficient Eq. (7–7), remains similar to the
correlogram. Although there are double summations in the correntropy coefficient, the computational complexity can be reduced to $O(N \log N)$ using the Fast-Gauss transform [123]. However, the ”narrowed” autocorrelation function increases computational complexity by including more delay terms.

7.3 Experiments

In this section, we present three experiments to validate our method. In the first two simulations, we compare our method with the conventional autocorrelation function [119], the third order cumulants function [116], and the narrowed autocorrelation function [103] in determining pitches for a single speaker and two combined speakers uttering different vowels. The synthetic vowels are produced by Slaney’s Auditory Toolbox [124]. For a fair comparison, we did not apply any post-processing on the correlogram as was used in [119]. The conventional autocorrelation function Eq. (7–1), autocorrelation of third order cumulants functions Eq. (7–4), narrowed autocorrelation functions Eq. (7–3)
Figure 7-8. Third order cumulants (top) and summary (bottom) for a mixture of vowels /a/ and /u/.

and correntropy functions Eq. (7–6) are presented after the same cochlear model. In the third experiment, the proposed method is tested using Bagshaw’s database which is a benchmark for testing PDAs [125].

7.3.1 Single Pitch Determination

Fig. 7-3 to Fig. 7-6 present the pitch determination results for a single synthetic vowel /a/ with fundamental frequency at 100Hz. The upper plots are the images of correlation functions, autocorrelations of third order cumulants, narrowed autocorrelations, and correntropy functions after the same cochlear model respectively. The bottom figures are the summaries of those four images. The kernel size $\sigma$ in the Gaussian kernel has been chosen to be 0.01 (we will discuss further kernel size selection in Sec. 8.3) and $L = 10$ in the narrowed autocorrelation function Eq. (7–3). The conventional autocorrelation, third order cumulants and narrowed autocorrelation are all able to produce peaks at 10ms corresponding to the pitch of the vowel. But they also generate
other erroneous peaks which might confuse pitch determination. On the contrary, the summary of correntropy-gram provides only one single and narrow peak at 10ms which is the pitch period of the vowel. And the peak is much narrower than those obtained from other methods. The correntropy-gram clearly shows a single narrow stripe across all the frequency channels which concentrates most of the energy. It is the indication of the fundamental frequency.

The fine structure of hyperbolic contours can also be clearly seen in the correntropy-gram. Its power spectrum energy spreads equally at all the harmonics. The rows of white spots across the correntropy-gram reflect the periodic structure. Particularly, the second harmonic shows two peaks during the time interval when the fundamental frequency exhibits one. This fine structure is the result of correntropy’s ability to contain all the harmonics of signal. However, the correlogram yields a much wider spread of energy across the frequency channels. The image with the autocorrelations of the third order cumulants
fail to present such structures. Although, the image of the narrowed autocorrelation functions is able to show some hyperbolic contours, the white vertical stripe at the fundamental frequency is much wider than that of the correntropy-gram and there are other white spots in the high frequency channels. These result in the wide peak at 10ms and other erroneous peaks in the summary of the narrowed autocorrelation functions. Our proposed method clearly outperforms the conventional autocorrelation function, third order cumulants method and narrowed autocorrelation function in single pitch determination case.

7.3.2 Double Pitches Determination

In this example, we consider pitch determination for a mixture of two concurrent synthetic vowels with /a/ ($F_0 = 100$ Hz) and /u/ ($F_0 = 126$ Hz) which are separated by four semitones. We compare the same four methods as the experiment above to
Figure 7.11. The ROC curves for the four PDAs based on correntropy-gram, autocorrelation, narrowed autocorrelation ($L = 15$), and autocorrelation of 3rd order cumulants in double vowels segregation experiment.

demonstrate that the correntropy function is able to determine two pitches presented in the mixture of two vowels.

Fig. 7-7 to Fig. 7-10 present the simulation results. The correlogram method result shown in Fig. 7-7 only shows one peak corresponding to the pitch of the vowel /a/ while no indication of the other vowel /u/ at time of 7.9ms is provided. The summary of correlogram resembles that of single vowel case in Fig. 7-3. The third order cumulants method in Fig. 7-8 fails to detect two pitches in the mixture signal. Although there are two small peaks at 10ms and 7.9ms which correspond to the two pitch periods respectively, their amplitudes are not large enough to be reliably detected. In Fig. 7-9, the summary of narrowed autocorrelation functions with $L = 15$ is able to produce only one peak at 10ms corresponding to the pitch period of vowel /a/, but there is no peak at 7.9ms. There are white spots in the low frequency channels in the image of narrowed autocorrelation functions which are the indications of the second vowel /u/. However, the amplitude is too small compared with that of vowel /a/ and the information is lost in the summary plot. A complex neural network oscillator has been used to separate the
Figure 7-12. The percentage performance of correctly determining pitches for both vowels for proposed PDA based on correntropy function and a CASA model.

channels dominated by different voices, and the summaries of individual channels are able to produce peaks corresponding to different vowels [101].

On the other hand, our method is able to detect two pitches from the mixture of two vowels. The kernel size $\sigma$ is set to 0.07 in this experiment. The correntropy-gram in Fig. 7-10 shows a white narrow stripe across high frequency channels at 10ms corresponding to the pitch period of the vowel /a/. These channels have center frequencies close to the three formant frequencies of vowel /a/ ($F_1 = 730\text{Hz}, F_2 = 1090\text{Hz}, F_3 = 2440\text{Hz}$). The hyperbolic structure can still be seen in the high frequency channels, but the lower frequency channels have been altered by the presence of vowel /u/. The three high energy white spots appear along the frequency channels centered at 300Hz which is the first formant of vowel /u/. The second white spot locates at 7.9ms matches the pitch period of vowel /u/. In the summary of correntropy-gram, the first peak at 10ms corresponds to the pitch period of vowel /a/. It is as narrow as the one in the single vowel case in Fig. 7-6. The second peak appears at 8.2ms which is only 4Hz off the true pitch frequency (126Hz). It is much less than the 20% gross error pitch determination evaluation criterion [126] or 10Hz gross error [99]. The second peak is also much wider than the one at 10ms. The
amplitude for the peak at 8.2ms is also smaller than that of peak at 10ms since the energy ratio is 5.2 times higher for vowel /a/. The pitch shift and peak broadening phenomenon is due to the fact that vowel /a/ dominates the mixture signal and it generates spurious peaks which blur that of vowel /u/. However, it is remarkable that our method, with the proper kernel size, is able to detect two pitches while all other algorithms fail in this experiment. This simulation clearly demonstrates the superiority of our method over the conventional correlogram, third order cumulants and narrowed correlation approaches for multipitch determination.

7.3.3 Double Vowels Segregation

To further investigate the performance of the proposed PDA, we generate a set of three vowels: /a/, /u/ and /i/ using Slaney’s Auditory Toolbox. Each vowel is synthesized on 5 pitches corresponding to differences of 0.25, 0.5, 1, 2 and 4 semitones from 100Hz, and the duration is 1s each. For every mixture of double vowels, one is always with the fundamental frequency at 100Hz, and the other constituent can be any vowel at any pitch value. In total, we have 45 mixtures of different combinations of vowels with different pitch values (3 vowels × 3 vowels × 5 pitches). The detection functions from each of four methods above have been normalized to 0 and 1. A threshold varies
between 0 and 1 to decide the peaks. If the difference between the detected pitch and reference is within a certain tolerance, the right pitch is detected. Since the minimum distance in this experiment is 0.25 semitone from 100Hz, which is 1.45 Hz, we select the tolerance to be 0.725Hz. Fig. 7-11 plots the ROC curves for the four pitch determination algorithms based on correntropy function, autocorrelation, narrowed autocorrelation and autocorrelation of third order cumulants. It clearly shows that our method outperforms the other three in double-vowel pitch detection. However, none is able to get 100% detection. Notice that the ROC curve for correntropy function contains many points of zero probability of false alarm, up to 45% of correct detection. This is due to the fact that correntropy function is able to suppress other erroneous peaks which are away from pitch positions and concentrate energy around fundamental frequencies. The performance of autocorrelation and third order cumulants are below 50% detection rate, irrespective of the number of false alarms generated, which means that most often the second largest peak is an harmonic of the highest pitch. This is not surprising since both functions fail to present two peaks for most mixtures in this experiment.

We also present the vowel identification performance to examine the discriminating ability of correntropy function at different semitones for the mixture of double vowels. In the experiment, the threshold is chosen such that the first two peaks are detected. We compare our results with a computational auditory scene analysis (CASA) model with a network of neural oscillators [101] in Fig. 7-12. The CASA model outperforms our method at 0.25, 0.5 and 0.5 semitones of F0 differences since it uses a sophisticated network of neural oscillators to assign different channels from ERB filter bank outputs to different vowels. Our method is just based on the simple summary function of correntropy-gram. The closer the two fundamental frequencies of two vowel become, the harder for correntropy function to produce two distinct peaks corresponding to different pitches. However, our method obtains comparable results to CASA model at 2 and 4 semitones of F0 differences. It suggests that our simple model is able to produce similar
results for double vowel segregation of 2 and 4 semitones of $F_0$ differences compared to the sophisticated CASA model. This certainly shows our technique is very promising.

### 7.3.4 Benchmark Database Test

We test our pitch determination algorithm with Bagshaw’s database \cite{125}. It contains 7298 males and 16948 females speech samples. The groundtruth pitch is estimated at reference points based on laryngograph data. These estimates are assumed to be equal to the perceived pitch. The signal is segmented into 38.4ms duration centered at the reference points in order to make the comparisons between different PDAs fair. The sampling frequency is 20kHz. The kernel size is selected according to Silverman’s rule for different segments. We use equation Eq. (7–7) to calculate the normalized correntropy functions such that the summary correntropy function at zero lag is unit. Since the pitch range for male speaker is 50-250 Hz and 120-400 Hz for female speaker, the PDA searches local maximums from 2.5 ms above in the summary correntropy function. We set the threshold to be 0.3 by trial and error so that every local maximum which exceeds 0.3 will be detected as a pitch candidate.

Table 7-1 summarizes the performance of various PDAs which are taken from \cite{126}, \cite{127}. The performance criterion is the relative number of gross error. A gross error occurs
Table 7-1. Gross error percentage of PDAs evaluation

<table>
<thead>
<tr>
<th></th>
<th>Male</th>
<th></th>
<th></th>
<th>Female</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>High (%)</td>
<td>Low (%)</td>
<td>High (%)</td>
<td>Low (%)</td>
<td>Weighted Mean (%)</td>
<td></td>
</tr>
<tr>
<td>PDA</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HPS</td>
<td>5.34</td>
<td>28.20</td>
<td>0.46</td>
<td>1.61</td>
<td>11.54</td>
<td></td>
</tr>
<tr>
<td>SRPD</td>
<td>0.62</td>
<td>2.01</td>
<td>0.39</td>
<td>5.56</td>
<td>4.95</td>
<td></td>
</tr>
<tr>
<td>CPD</td>
<td>4.09</td>
<td>0.64</td>
<td>0.61</td>
<td>3.97</td>
<td>4.63</td>
<td></td>
</tr>
<tr>
<td>FBPT</td>
<td>1.27</td>
<td>0.64</td>
<td>0.60</td>
<td>3.35</td>
<td>3.48</td>
<td></td>
</tr>
<tr>
<td>IPTA</td>
<td>1.40</td>
<td>0.83</td>
<td>0.53</td>
<td>3.12</td>
<td>3.22</td>
<td></td>
</tr>
<tr>
<td>PP</td>
<td>0.22</td>
<td>1.74</td>
<td>0.26</td>
<td>3.20</td>
<td>3.01</td>
<td></td>
</tr>
<tr>
<td>SHR</td>
<td>1.29</td>
<td>0.78</td>
<td>0.75</td>
<td>1.69</td>
<td>2.33</td>
<td></td>
</tr>
<tr>
<td>SHAPE</td>
<td>0.95</td>
<td>0.48</td>
<td>1.14</td>
<td>0.47</td>
<td>1.55</td>
<td></td>
</tr>
<tr>
<td>eSRPD</td>
<td>0.90</td>
<td>0.56</td>
<td>0.43</td>
<td>0.23</td>
<td>0.90</td>
<td></td>
</tr>
<tr>
<td>Correntropy</td>
<td>0.71</td>
<td>0.42</td>
<td>0.35</td>
<td>0.18</td>
<td>0.71</td>
<td></td>
</tr>
</tbody>
</table>

when the estimated fundamental frequency is more than 20% off the true pitch value.

The percent gross errors by gender and by lower or higher pitch estimates with respect to the reference are given in Table 7-1. The weighted gross error is calculated by taking into account the number of pitch samples for each gender. It clearly shows that for this particular database correntropy based PDA outperforms others.

7.4 Discussions

In all the previous experiments, a specific kernel size $\sigma$ in the Gaussian kernel Eq. (3-3) needs to be selected by the user (free parameter). Actually, the kernel size plays an important role in the performance of our method since it determines the scale at which the similarity is going to be measured. It has been shown that kernel size controls the metric of the transformed signal in the RKHS [65]. If the kernel size is set too large, the correntropy function approaches the conventional correlation function and fails to detect any nonlinearity and higher order statistics intrinsic to the data; on the other hand, if the kernel size is too small, the correntropy function loses its discrimination ability. One practical way to select the kernel size is given by the Silverman’s rule [120]

$$\sigma = 0.9AN^{-1/5},$$
where $A$ is the smaller value between standard deviation of data samples and data interquartile range scaled by 1.34, and $N$ is the number of data samples.

To illustrate the effect of different kernel sizes, we simulate the summary of correntropy functions for the same experiments setup in Sec. 8.2 with different kernel sizes in Fig. 7-13 and Fig. 7-14. It can be seen that if the kernel size is large, $\sigma = 1$ here, the summaries of correntropy functions approach those of correlation functions shown in Fig. 7-3 and Fig. 7-7. As kernel size approaches the kernel size given by Silverman’s rule, $\sigma = 0.01$ for the single vowel /a/ case and $\sigma = 0.07$ for a mixture of /a/ and /u/, the summary of correntropy functions starts to present a large and narrow peak corresponding to the pitch of vowel /a/ and show the other vowel /u/. If the kernel size is too small, $\sigma = 0.001$ here, the summary of correntropy functions loses its ability to present two vowels. This is shown in the bottom plot of Fig. 7-14.

7.5 Conclusion

A novel pitch determination algorithm is proposed based on the correntropy function. The pitch estimator computes the correntropy functions for each channel of an ERB filter bank, and adds across all the channels. Simulations on single and double vowel cases show that the proposed method exhibits much better resolution than the conventional correlation function, third order cumulants method and narrowed correlation function in single and double pitches determination. This suggests that correntropy can discriminate better pitch when two different speakers speak in the same microphone. This is essential in computational auditory scene analysis (CASA). Although these results are preliminary and much further work is needed to evaluate the methods, this technique seems promising for CASA. The automatic selection of the kernel size or of a multiple kernel size analysis need to be further investigated to automate the pitch determination algorithm. The future work also includes incorporating correntropy-gram channel selection to enhance the discriminating ability of proposed method in multiple pitches determination. A benchmark database test for various PDAs shows that the proposed PDA outperforms some others
with available results for this dataset. Since correntropy creates many different harmonics of each resonance present in the original time series due to the nonlinearity, it may also be useful for perceptual pitch determination.
CHAPTER 8
CORRENTROPY COEFFICIENT AS A NOVEL SIMILARITY MEASURE

In this chapter, we apply the proposed correntropy coefficient (4–27) to characterize the similarity between multi-channel signals. Preliminary experiments with simulated data and multichannel electroencephalogram (EEG) signals during behavior studies elucidate the performance of the new measure versus the well established correlation coefficient.

8.1 Introduction

Quantification of dynamical interdependence in multi-dimensional complex systems with spatial extent provides a very useful insight into their spatio-temporal organization. In practice, the underlying system dynamics are not accessible directly. Only the observed time series can help decide whether two time series collected from the system are statistically independent or not and further elucidate any hidden relationship between them. Extracting such information becomes more difficult if the underlying dynamical system is nonlinear or the couplings among the subsystems are nonlinear and non-stationary.

There has been extensive research aimed at detecting the underlying relationships in multi-dimensional dynamical systems. The classical methodology employs a linear approach, in particular, the cross correlation and coherence analysis [128]. Cross correlation measures the linear correlation between two signals in the time domain, while the coherence function specifies the linear associations in the frequency domain by the ratio of squares of cross spectral densities divided by the products of two auto-spectra. There have been several extensions of correlation to more than two pairs of time series such as directed coherence, directed transfer functions and partial directed coherence [129]. Unfortunately, linear methods only capture linear relationships between the time series, and might fail to detect nonlinear interdependencies between the underlying dynamical subsystems.
Nonlinear measures include mutual information and state-space methods. One technique is the generalized mutual information function [130]. However, a large quantity of noise-free stationary data is required to estimate these measures based on information theory, which restricts their applications in practice. Another method is the phase synchronization where the instantaneous phase using Hilbert transforms is computed and interdependence is specified in terms of time-dependent phase locking [131]. The state-space methodologies include similarity-index and synchronization likelihood. The similarity-index technique and its modifications compute the ratio of average distances between index points, their nearest neighbors and their mutual nearest ones [132, 133]. Stam et al. proposed the synchronization likelihood to offer a straightforward normalized estimate of the dynamical coupling between interacting systems [134]. There are several drawbacks associated with these techniques based on state space embedding. Estimating the embedding dimension of times series corrupted by measurement noise for a valid reconstruction, searching a suitable neighborhood size and finding a constant number of nearest neighbors are a few of many constraints that severely affect the estimation accuracy.

In this chapter, we use the correntropy coefficient Eq. (4–27) as a novel similarity measure to quantify the inter-dependencies among multi-channel signals. In practice, the estimate of correntropy coefficient Eq. (4–28) is calculated between two time series of interests to measure the similarity.

8.2 Experiments

We test the correntropy coefficient on simulated data set and real world EEG signals in the experiments.

8.2.1 Two Unidirectionally Coupled Hénon maps

First, we test two unidirectionally coupled Hénon maps which has been extensively used to validate many synchronization measure in the literature [133, 134]. We apply the correntropy coefficient in detecting nonlinear interdependence of two unidirectionally
coupled Hénon maps defined as

\begin{align}
x_1(n+1) &= 1.4 - x_1^2(n) + b_x x_2(n) \quad (8-1) \\
x_2(n+1) &= x_1(n) \quad (8-2)
\end{align}

for the driver, represented as system X, and

\begin{align}
y_1(n+1) &= 1.4 - [C x_1(n) + (1 - C) y_1(n)] y_1(n) + b_y y_2(n) \quad (8-3) \\
y_2(n+1) &= y_1(n) \quad (8-4)
\end{align}

for the response, represented as system Y. System X drives system Y with nonlinear coupling strength \(C\). \(C\) ranges from 0 to 1 with 0 being no coupling and 1 being complete coupling. Parameters \(b_x\) and \(b_y\) are both set to 0.3 as canonical values for the Hénon map when analyzing identical systems, and to 0.3 and 0.1 respectively for nonidentical ones.

For each coupling strength, we discard the first 10000 iterated time series as transient and obtain the next 500 data points for experiments. The correntropy coefficient \(\eta\) is calculated between the first component of system X, \(x_1\), and the first component of system Y, \(y_1\).

In the following simulations, we aim to address these questions: (1) whether correntropy coefficient increases when the coupling strength \(C\) between X and Y for both identical and nonidentical systems increases; (2) How robust is correntropy coefficient to different level measurement noises in driver, response and both systems? (3)How sensitive is correntropy coefficient to time dependent sudden change in the dynamics of interacting systems due to coupling strength? (4) Can correntropy coefficient detect nonlinear coupling between driver and response? (5) How is correntropy coefficient affected by kernel size and data length?

### 8.2.1.1 Variation of Correntropy Coefficient with Coupling Strength

First in Fig. 8-1, we plot the averaged correntropy coefficient \(\eta\) as a function of coupling strength \(C\) for identical map \((b_x = b_y = 0.3)\) and nonidentical map \((b_x = 0.3)\)
and $b_y = 0.1$) over 10 realizations of different initial conditions. The error bars denote the standard deviation over the different realizations. Fig. 8-1(a) shows the identical map where the kernel size used in Gaussian kernel has been chosen to be 0.001 according to the Silverman’s rule Eq. (8–5). For identical chaotic systems, perfect synchronization can be generated with sufficient degrees of coupling [135]. This can be seen from the fact that the correntropy coefficient $\eta = 1$ for $C \geq 0.7$ in Fig. 8-1(a) indicating perfect synchronization occurs between two coupled systems. The critical threshold $C = 0.7$ corresponds to the point when the maximum Lyapunov exponent of the response system becomes negative and identical synchronization between the systems takes place [133]. On the other hand, the correntropy coefficient $\eta = 0$ for $C < 0.7$ suggesting no synchronization even though two systems are weakly coupled. Similar results have been reported using other nonlinear interdependence measurement in [133, 136, 137].

Fig. 8-1(b) shows the result for unidirectionally coupled nonidentical systems ($b_x = 0.3, b_y = 0.1$). The kernel size is set to 0.4. In this case, identical synchronization is not possible and the driver has higher dimension than the response. The sharp increase of the correntropy coefficient at point $C = 0.7$ as in the identical synchronization situation can not be observed here. But the correntropy coefficient shows a consistent monotonic
Figure 8-2. Influence of different noise levels on correntropy coefficient.

increase with respect to coupling strength except for the region $0.1 < C < 0.3$. The local hump in this region can also be seen in [133, 134, 137]. It is suggested that the cause is most likely due to the local minimum of the maximum Lyapunov exponent [133, 138] in this region.

8.2.1.2 Robustness Against Measurement Noise

A good interdependence measure should be robust against noise. Next, we analyze the robustness of correntropy coefficient when time series are contaminated with noise. There are two types of noise in general: intrinsic noise and additive measurement noise. We only consider measurement noise here which does not perturb the inherent dynamics of systems. Independent realizations of white noise are added to driver, response and both systems separately. The signal-to-noise (SNR) ratio is set to be 10dB and 1dB respectively to test the performance of correntropy coefficient at different noise intensity. 500 data samples are used to calculated the correntropy coefficient, averaged over 20 realizations. Fig. 8-2 plots the correntropy coefficient for unidirectionally coupled identical Hénon map ($b_x = b_y = 0.3$) with white noise in response, driver and both systems. Kernel size is chosen to be 0.04 for SNR = 10dB and 0.08 for SNR = 1dB respectively. Notice that the correntropy coefficient curves with noise become less smooth than that of noise-free
one, but the sharp increase at $C = 0.7$ is still obvious for both noise intensities. When noise level is high (SNR = 1dB), the correntropy coefficient curve is more zigzag than that of 10dB case, however it can still detect increases in the coupling strength. The figure also suggests that whether noise is added into driver, response or both system, the performance of correntropy coefficient is very similar. Fig. 8-3 presents the results for non-identical Hénon map ($b_x = 0.3$ and $b_y = 0.1$) with white noise in response, driver and both systems. Kernel size is selected to 0.05 for SNR = 10dB and 0.2 for SNR = 1dB respectively. The values of the correntropy coefficients at different coupling strength are averaged over 20 independent realizations. In both levels of noise case, the correntropy coefficients consistently increase with respect to coupling strength. Also the effect of noise in response, driver or both systems does not make big differences. Notice that the local hump around the region $0.2 < C < 0.4$ is still observable for all cases. These results show that the correntropy coefficient is fairly robust even in the case of considerably noisy data.

8.2.1.3 Sensitivity to Time-dependent Dynamical Changes

Next we test how sensitive the correntropy coefficient is to time dependent sudden change in the dynamics of interacting systems due to coupling strength. In experiment, change in coupling strength can cause sudden change in the dynamics of interacting
systems, which basically generates non-stationarity in time series. To study such transient
dynamical phenomenon, both identical \((b_x = b_y = 0.3)\) and non-identical \((b_x = 0.3\) and \(b_y = 0.1\)) Hénon maps are considered here. Dynamical systems are coupled only during
a single epoch and otherwise uncoupled for both cases \([134, 137]\). We set the coupling
strength \(C = 0\) for \(n \leq 10150\) and \(n \geq 10250\) and \(C = 0.8\) for \(10150 < n < 10250\).
Only 400 data samples are plotted after the first 10000 data are discarded as transient.
The sliding window used to compute the correntropy coefficient is chosen to contain
8 data. Kernel size is set to 0.2 for identical map and 0.3 for non-identical map. The
results are averaged over 20 independent realizations of different initial conditions ranging
0 to 1. Fig. 8-4 plots the correntropy coefficient curves for identical and non-identical
maps. In uncoupled regions, \(\eta\) fluctuates around 0.01 baseline for identical map and
0.001 for non-identical map. A sharp and clear increase occurs at \(t = 150\) when 0.8
coupling strength between systems \(\mathbf{X}\) and \(\mathbf{Y}\) is introduced, and there is a sharp and
clear decrease in \(\eta\) falling off back to the baseline level when coupling strength between
two systems reduces to zero at \(t = 250\). The interval where \(\eta\) is noticeably higher than
the baseline level matches nicely to the coupling interval. This phenomenon is observed
both in identical and non-identical Hénon maps. Therefore, the correntropy coefficient

Figure 8-4. Time dependent of correntropy coefficient.
is potentially able to detect sudden change in the coupling between two interacting dynamical systems with a high temporal resolution, which makes this measure suitable for non-stationary data sets.

### 8.2.1.4 Effect of Kernel Width

We have discussed the importance of the kernel width in the performance of the correntropy coefficient because it is a parametric measure in previous section. Here we demonstrate this on unidirectionally coupled identical and non-identical Hénon maps. Fig. 8-5 shows the correntropy coefficient curves of different kernel width for unidirectionally coupled identical Hénon map. When the kernel width is chosen too large, $\sigma = 0.1, 0.5, 1$, in this case, correntropy coefficient produces erroneous results in the unsynchronized region $0 < C < 0.7$. The results for non-identical Hénon map are presented in Fig. 8-6. It can be seen that if kernel width is too small, the increase of the correntropy coefficient with respect to the coupling strength is not as obvious as those of suitable kernel width ($\sigma = 0.4$ here). While the kernel width is too larger, the results of the correntropy coefficient approach to those of conventional correlation coefficient. In both figures, we see that the correntropy coefficients can either increase or decrease as the kernel width increases. These observations are consistent with our theoretical analysis in previous section.
8.2.1.5 Ability to Quantify Nonlinear Coupling

To demonstrate that correntropy coefficient $\eta$ is able to detect the nonlinear coupling between systems $X$ and $Y$, we first compare our measure with the conventional correlation coefficient and one of the well-known nonlinear synchronization measure, similarity index in [133], on identical Hénon maps. Fig. 8-7 shows the correlation coefficient, the correntropy coefficient and the similarity index as functions of coupling strength $C$.

The correntropy coefficient generates exactly the same result as the similarity index. On the other hand the conventional correlation coefficient performs erratically in the unsynchronized region $C < 0.7$. This clearly demonstrates that the correntropy coefficient outperforms the correlation coefficient in characterization of nonlinear coupling between two dynamical systems. Compared to the similarity index, the correntropy coefficient has the advantage of avoiding estimating embedding dimension, choosing nearest neighborhood and other problems associated with state space embedding method [133, 139]. The computational complexity of the correntropy coefficient is still manageable, and the kernel size is easy to estimate.

We also use multivariate surrogate data to further investigate that the correntropy coefficient is sensitive to nonlinear coupling. Prichard et al. introduced surrogate method
in [140]. This method has been applied to detect nonlinear structure in time series.

Basically, to generate multivariate surrogate data, first the Fourier transform is applied to each of time series, then a common random number is added to each of the phases and an inverse Fourier transform is applied. The resulting time series have identical power spectra and cross power spectra as the original time series, but any nonlinear coupling among the time series has been destroyed. In simulation, we use TISEAN package [141] to generate 19 realizations of the surrogate data for the time series \( x_1(t) \) in Eq. (8–1) and \( y_1(t) \) in Eq. (8–3) for each different coupling strength for the unidirectionally coupled non-identical Hénon map. Then we compute the correntropy coefficient for both the original and the surrogate data with respect to different coupling strength. Fig. 8-8 plots the correntropy coefficient curve for the original data and the mean value of 19 correntropy coefficients for the surrogate data with the corresponding maximal and minimal values as error bars.

To quantify the significance level, we calculate the Z-Score as

\[
Z = \frac{|v_{\text{orig}} - \mu_{\text{surr}}|}{\sigma_{\text{surr}}}
\]

where \( v_{\text{orig}} \) is the correntropy coefficient value for the original data, \( \mu_{\text{surr}} \) and \( \sigma_{\text{surr}} \) are the mean and the standard deviation for the surrogate data respectively. Table 8-1 presents the Z-Score values for different coupling strength. With the exception of \( C = 0.2 \) and 0.4, the Z-Score values are significantly larger than 1.96 which means the nonlinear coupling

Figure 8-7. Comparison of correlation coefficient, correntropy coefficient and similarity index.
has been detected with a probability $p < 0.05$. These results clearly demonstrate that the correntropy coefficient is sensitive to the nonlinearity of the dependence between two coupled systems.

### 8.2.2 EEG Signals

In the second experiment, we applied the correntropy coefficient to real EEG signals. The electrical potentials on the surface of the scalp of a human subject were measured and recorded with the NeuroScan EEG system (NeuroScan Inc., Compumedics, Abbotsford, Australia). A 64-channel cap was used with electrode locations according to the extended international 10/20 system and with a linked-earlobe reference. Horizontal and vertical electrooculogram (HEOG and VEOG) signals were also recorded for artifact rejection using two sets of bipolar electrodes. The data sampling rate was fixed at 1000Hz and the online band-pass filter range was set to be maximally wide between 0.05Hz and 200Hz. Subjects were presented repeatedly (200 times) with uni-modal auditory and visual stimuli delivered in the central visual and auditory spaces simultaneously and with the same strength to the left/right eyes and ears, as well as with simultaneous cross-modal combinations. For the purpose of this study, only the uni-modal data was used. The visual
stimuli consisted of 5x5 black and white checkerboards presented for 10ms, while the auditory stimuli were 2000Hz tones with durations of 30ms. The time interval between the stimuli in any of the experimental conditions was random between 1500ms and 2000ms. Following standard eye-movement artifact rejection procedures and segmentation into single epochs with alignment at the onset of the stimuli, all artifact-free epochs were averaged and normalized to zero mean and unit variance and low-pass filtered at 0-40Hz for further analysis. We then applied the correntropy coefficient to the averaged data to quantify the bilateral synchronization or couplings among the corresponding sensory areas of the brain. In order to test whether the correntropy coefficient was able to detect any nonlinear couplings in the EEG signals, the results were compared to the conventional correlation coefficient. A window size of 20ms data is used to calculate both measures corresponding to the duration of a single dipole activation in the cortex [142]. The kernel width $\sigma$ in Gaussian kernel used in correntropy coefficient was chosen to be 0.4.

Fig. 8-9 (a) and (b) show plots of the correlation and correntropy coefficients for the auditory areas of the brain as a function of time after the subject was exposed only to the audio stimuli. Several bilaterally-symmetrical pairs of electrodes were selected in the vicinity of the auditory cortex, so that both measures were computed for pairs FC5-FC6, FC3-FC4, C5-C6, C3-C4, CP5-CP6, CP3-CP4. As shown in Fig. 8-9 (a) and (b), there are two distinct time intervals 0-270ms and 270-450ms in the auditory response. Both correlation and correntropy coefficients drop at 270ms. This suggests that both measures are able to detect the changes in inter-hemispheric synchronization of the auditory regions. However, as the electrodes are chosen in different locations away from the auditory cortex, it is expected that during the synchronization phase (0-270ms) the synchronization measures for different pairs should be different. Fig. 8-9 (a) shows that

<table>
<thead>
<tr>
<th>C</th>
<th>0</th>
<th>0.1</th>
<th>0.2</th>
<th>0.3</th>
<th>0.4</th>
<th>0.5</th>
<th>0.6</th>
<th>0.7</th>
<th>0.8</th>
<th>0.9</th>
<th>1</th>
</tr>
</thead>
</table>
the correlation coefficients for all 6 pairs are grouped together and are unable to detect the difference in activation, while Fig. 8-9 (b) suggests that the correntropy coefficient can differentiate successfully the synchronization strength among different areas of the cortex above the left and right auditory regions. Notably, as expected from previous studies, pairs FC5-FC6 and FC3-FC4 exhibit stronger synchronization strength than the others, while most posterior pairs CP5-CP6 and C5-C6 have weaker synchronization strength. Also the synchronization patterns reveal lateral similarity in time for the pairs FC5-FC6 and FC3-FC4, for CP5-CP6 and C5-C6, and for CP3-CP4 and C3-C4. Furthermore the correntropy coefficients for pairs C5-C6, C3-C4 and CP3-CP4 peak simultaneously at 90ms which corresponds to the first mean global field power (MGFP) peak of the EEG signal.
Figure 8-10. Comparison of correlation coefficient and correntropy coefficient in characterization of synchronization among occipital cortex for visual stimulus EEG signal.

These differences indicate that the correntropy coefficient is more sensitive and is able to extract more information as a synchronization measure than the conventional correlation coefficient.

We also compared both measures when applied to the visual cortical areas. The measures are presented in Fig. 8-10 as a function of time when the subject is exposed only to visual stimuli. Again, a window size of 20ms data is used to compute both the correlation and the correntropy coefficients, and the kernel width $\sigma$ is again set to 0.4 as in the previous case. We also chose bilaterally symmetrical pairs of electrodes O1-O2, PO7-PO8, PO5-PO6, P7-P8, P5-P6 and P3-P4. In Fig. 8-10 (b) the correntropy coefficients for all pairs except for O1-O2 show similar synchronization patterns. The
correntropy coefficient increases at first, then reaches a peak around 275ms, after which it drops to lower levels. The maximum values of the correntropy coefficients around 275ms correspond to the peak P1 in the visual evoked potential [143]. As expected the synchronization between occipital channels O1 and O2 has the maximum strength and stays high until it decreases around 350ms. Thus the correntropy coefficient shows that the extra-striate visual networks become increasingly recruited and synchronized until about 275ms after the stimulus onset, while the primary visual cortex is highly synchronous for a longer period of time, until about 350ms after onset. The channels pair P7 and P8 exhibits the weakest synchronization strength since it is located the farthest away from the primary visual cortex compared to other electrode pairs. On the other hand, the correlation coefficients for most channel pairs group together and display the same level of synchronization until its sharp decrease at around 500ms (except for P7-P8). The synchronization between P7 and P8 has irregular patterns with a local minimum around 200ms. This comparison clearly demonstrates that also in this case the correntropy coefficient measure outperforms the correlation coefficient in the quantification of the EEG signal coupling between the bilateral occipital regions of the brain in response to visual stimuli.

8.3 Discussions

In this section, we explore more details about the correntropy coefficient both in theoretical analysis and practical implementation such as the geometrical interpretation, connections to information theory, the important role of the kernel size, and the scaling factor of time series.

8.3.1 Kernel Width

The variance of the Gaussian kernel function is normally called kernel width or kernel size. The kernel size should be considered a scale parameter controlling the metric in the projected space. From the geometrical perspective, the kernel size decides the length of each of the nonlinearily transformed vectors and the angle between them in the RKHS
Because $\|\Psi(x)\| = \sqrt{\kappa(0) - \frac{1}{N^2} \sum_{i=1}^{N} \sum_{j=1}^{N} \kappa(x_i, x_j)}$. It can be easily seen that the vector length monotonically decreases as the kernel width increases, while the centered crosscorrentropy exhibits more complex pattern since it also depends on the nature of the relationship between two random variables. The kernel width controls the ability of the centered correntropy to capture the nonlinear couplings intrinsic in the signals. If the kernel width is too large, the correntropy coefficient loses the ability to detect the nonlinearity and approaches the conventional correlation coefficient; when the kernel width is too small, the nonlinear transformations $\Phi$ in Eq. (4–3) and $\Psi$ in Eq. (4–29) cannot interpolate between data points. This can also be verified to apply the Taylor series expansion to the Gaussian kernel where the kernel width appears as the weighting parameter in both second and higher order moments. The effect of the kernel size on different order moments is scaled by the power up to $2^k$ where $k$ is the moment order.

When the kernel size is too large, the contribution of higher order statistics decays rapidly and the centered crosscorrentropy approaches the conventional crosscovariance function; on the other hand, when the kernel size is too small, the effect of higher order moments overweighs the second order one. An appropriate kernel size should maintain the balance of second order and higher order statistics of the signal.

Therefore a good choice of the kernel parameter is crucial for obtaining the good performance of the proposed method. There are two ways of handling the selection of kernel size. One is to seek an optimal kernel size. The cross-validation has been one of the mostly used methods in machine learning field to choose an appropriate kernel width. Other approaches include the Silverman’s rule of thumb which is given by [120]:

$$\sigma = 0.94A N^{-1/5}, \quad (8–5)$$

where $A$ is the smaller value between standard deviation of data samples and data interquartile range scaled by 1.34, and $N$ is the number of data samples. The Silverman’s rule is easy to use to choose a good kernel size, hence we will set the kernel width
according to Eq. (8–5) throughout the paper. Alternatively, the kernel size can be thought as a scale parameter that provides different looks to the dependence among the variables. Just like in wavelets, the kernel size is able to analyze the dependencies at different resolutions. Since many real world signals are very complex, this multi-resolution analysis may elucidate better the relationships.

### 8.3.2 Scaling Effect

Because the nonlinear transformations $\Phi$ in Eq. (4–3) and $\Psi$ in Eq. (4–29) are nonlinear, any scaling of the original random variables results in different performance of the correntropy coefficient. Unlike the conventional correlation coefficient which is insensitive the amplitude scaling of the signals and only measures the similarity of the signals through the time, the correntropy coefficient measures both the time and amplitude similarities between two signals. Therefore, in certain applications, it is vital to normalize both signals before applying the correntropy coefficient. For example, the amplitudes of EEG signals are highly dependent on the different electrode impedances. It is important to normalize all channels of EEG signals to the same dynamical range.

### 8.4 Conclusion

In this chapter, we apply the correntropy coefficient as a novel nonlinear interdependence measure. Due to a positive definite kernel function, the correntropy coefficient implicitly maps the original random variables or time series into an infinite dimensional reproducing kernel Hilbert space which is uniquely induced by the centered crosscorrentropy function and essentially computes the cosine of the angle between the two transformed vectors. Orthogonality in RKHS $\mathcal{H}_U$ corresponds to independence between original random variables. Comparisons between the correntropy coefficient and the conventional correlation coefficient on simulated two unidirectionally coupled Hénon maps time series and EEG signals collected from sensory tasks clearly illustrate that the correntropy coefficient is able to extract more information than the correlation coefficient in quantification of synchronization between interacting dynamical systems.
CHAPTER 9
CONCLUSIONS AND FUTURE WORK

9.1 Conclusions

In this dissertation, we have analyzed the recently proposed correntropy function \[65\] and presented a new centered correntropy function from Time-domain and frequency-domain approaches. It is demonstrated that correntropy and centered correntropy functions not only capture the time and space structures of signals, but also partially characterize the high order statistical information and nonlinearity intrinsic to signals. Correntropy and centered correntropy functions have rich geometrical structures. Correntropy is positive definite and centered correntropy is non-negative definite, hence by Moore-Aronszajn theorem they uniquely induce reproducing kernel Hilbert spaces. Correntropy and centered correntropy functions combine the data dependent expectation operator and data independent kernels to form another data dependent operator. One perspective to work with correntropy and centered correntropy functions is to treat them as “generalized” correlation and covariance functions on nonlinarly transformed random signals via the data independent kernel functions. Those nonlinearly transformed signals appear on the sphere in the RKHS induced by the kernel functions if isotropic kernel functions are used. The other perspective is to directly work with the RKHS induced by the correntropy and centered correntropy functions directly. Now the nonlinearly transformed signals in the RKHS is no longer stochastic but rather deterministic. The RKHS induced by the correntropy and centered correntropy functions includes the expectation operator as embedded vectors. The two views further our understandings of correntropy and centered correntropy functions in geometrical perspective. The two reproducing kernel Hilbert spaces induced by kernel functions and correntropy functions respectively represent stochastic and deterministic functional analysis.

The correntropy dependence measure is proposed based on the correntropy coefficient as a novel statistical dependence measure. The new measure satisfies all the fundamental
desirable properties postulated by Renyi. We apply the correntropy concept in pitch
determination, and nonlinear component analysis. The correntropy coefficient is also
employed as a novel similarity measure to quantify the inter-dependencies of multi-channel
signals.

9.2 Future work

As we have provided a new methodology to explicitly construct a reproducing kernel
Hilbert space induced by the data independent kernel functions, we would like to apply
the same methodology to construct the RKHS induced by the data dependent correntropy
function directly. This shall allow us to analyze the functional basis that consists of the
RKHS. Moreover, if we can have an explicit expression for those functionals, then we can
develop algorithms that are not restricted to the inner product. We have only test our
correntropy PCA algorithms on artificial data sets. It is desired to apply the algorithms to
real data set. Some of the future work might includes

1. applying the parametric correntropy coefficient in detecting nonlinear coupling for
   EEG signal,
2. investigating the relationship between the higher order statistics and correntropy,
3. applying correntropy pitch determination algorithm in multiple pitches tracking.
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BIOGRAPHICAL SKETCH

Jianwu Xu was born in Wenzhou, China on December 7, 1979. He received his Bachelor of Engineering in Electrical Engineering from Zhejiang University, Hangzhou, China, in June 2002. Since October 2002, he has been working towards his Ph.D. in the Electrical and Computer Engineering Department at University of Florida, under the supervision of Dr. Jose Principe with support from Alumni Graduate Fellowship from University of Florida and NSF grant ECS-0601271. During the summer of 2006, he visited RIKEN Brain Science Institute in Tokyo, Japan and worked with Dr. Andrzej Cichocki on EEG synchronization in the Laboratory for Advanced Brain Signal Processing. His current research interests broadly include information theoretic learning, adaptive signal processing, control and machine learning. He is a member of IEEE, Tau Beta Pi and Eta Kappa Nu.