Testing independence in networks via family of network metrics

Abstract

Investigating how network structures are associated with nodal attributes of interest is a core problem in network science. As the network topology is a structured and often high-dimensional object, many traditional nonparametric tests are no longer applicable and parametric approaches are dominant in graph inferences. Here we propose a new procedure to testing dependence between graphs and attributes, via diffusion distances and distance-based correlation testing. We demonstrate that our nonparametric method not only yields a consistent test statistic under common network models, but also significantly surpasses the testing power of existing benchmarks under various circumstances.

Keywords: network dependence, distance correlation, exchangeable graph, diffusion maps

1 Introduction

Propelled by increasing demand and supply of graph data from various disciplines, the ubiquitous influence of network inferences has motivated numerous recent advances and applications in statistics, physics, computer science, biology, social science, etc., which further poses many new challenges to data scientists. One of the most fundamental statistical questions is to determine and characterize the relationship among multiple modalities of a given data set, for which the first step is to test the existence of any dependency. However, the lack of a principal notion of correlation in the graph domain has not only hindered the progress of nonparametric dependency testing methods, but also deterred a rich literature of statistical techniques in other inferences (e.g., regression, feature screening, two-sample test) from being directly applied to graphs.

Statisticians have long considered the problem of revealing the relationship between two data sets. The most classical approach is the Pearson's correlation [12], which determine the existence of linear relationship via a correlation coefficient in the range of [-1,1], with 0 indicating no linear association while ± 1 indicating perfect linear association. To capture all types of dependencies not limited to linear relationship, many new correlation measures and nonparametric statistics have been suggested to test independence between two random vectors [3-5, 10, 13, 14, 16, 18, 19]. In particular, the distance correlation by Szekely et al. [19] is the first correlation measure that is consistent against all possible dependencies (with finite moments), and the multiscale generalized correlation (MGC) statistic by Shen et al. [16] inherits the same consistency of distance correlation with significantly better finite-sample testing powers under high-dimensional and nonlinear dependencies, via defining a family of distance-based local correlations and efficiently searching the optimal correlation for testing.

Despite of many recent development, the network data, different from a random vector, still suffers from a dearth of proper analysis, mainly due to its structured and high-dimensional nature. Mathematically, a graph or, equivalently, network $\mathbf{G} = (V, E)$ consists of a set V of nodes (or vertices) together with a set E of edges, which is often represented via an adjacency matrix $\mathbf{A} = \{A_{ij} : i, j = 1, ..., n\}$, e.g. for an unweighted and undirected network, $A_{ij} = 1$ if node i and node j are connected by an edge, and zero otherwise. Therefore, \mathbf{A} is a symmetric square matrix that does not satisfy traditional data assumptions, e.g., each observation can be assumed independently and identically distributed, the sample size increases faster than the feature dimension, etc., which is a huge obstacle for directly applying conventional statistical methods.

When it comes to investigating the relationships between the data in network, a core problem is to detect dependency between network topology and nodal attributes, i.e., certain properties defined on nodes. For example, each person on Facebook having a number of different attributes, e.g., occupations, sex, personal behaviors, etc., are interacting each other via the social network; in neuro-science, each brain region has its own functionality, but is connected with other regions in the brain map. Identifying dependency between network and nodal attributes has primarily focused on their relationship explained only by network model under the boundary of model assumption [2, 7, 20]. Even though Fosdick and Hoff [2] suggested estimating node-specific network factors without restricting the dimensions, a fundamental difficulty of model-based independence tests is still lingering from the fact that not all networks exhibit the structures described by known network models. To our best knowledge so far, there is no principled method to compute a model-free correlation measure for testing network dependency.

To overcome the obstacles due to the distinct structure of network data and also due to the limitations of model-based method, we first define a family of distance metrics on network data via the diffusion maps, then apply nonparametric testing method of MGC utilizing the diffusion distance of the network topology and the Euclidean distance of the nodal attributes. Theoretical results show that under very mild condition, the diffusion maps, acting as a node-specific random vector, can allow distance-based correlation measures to be consistent in testing network dependencies. Moreover, the MGC statistic offers major power improvement under various scenarios in finite-sample testing. We further illustrate the advantages of diffusion maps and MGC over the existing benchmarks via comprehensive simulations under popular network models.

2 Results

2.1 Diffusion Maps and Diffusion Distances

In this section, we introduce a family of network geometries of an exchangeable graph [1] that can furnish conditional i.i.d. samples. A graph G is called exchangeable if and only if its adjacency matrix Ais jointly exchangeable [11], i.e. for every permutation σ of n, $(A_{ij}) \stackrel{d}{=} (A_{\sigma(i)\sigma(j)})$. Most statistical network models satisfy this property, including the Erdos-Renyi model, latent position model [2], stochastic block model [15], random dot product model [17], etc. [6, 8, 21, 22]. Coifman and Lafon [1] proposed multiscale geometries of data called diffusion maps, which inherit every local relation between the nodes when applied in network. The diffusion maps are constructed by iterating transition matrix which determines random walk on graph. To be specific, when given an $n \times n$ adjacency matrix A, we can define a $n \times n$ transition matrix of **P** where $P_{ij} = A_{ij} / \sum_{j=1}^{n} A_{ij}$ indicates the probability of moving forward from node i to node $j, i, j = 1, \dots, n$. In the so-called diffusion process, we basically run random walks by iterating this transition matrix, assuming that we are traveling from one node to the other once at each iteration with given transition probability. Then diffusion maps accordingly locate each node's position every diffusion time and provide node-specific multivariate coordinates. The diffusion maps at time t, $\mathbf{U}_t = \{\mathbf{U}_t(1), \dots, \mathbf{U}_t(n)\},$ can also be represented using a discrete set of real nonzero eigenvalues $\{\lambda_r\}$ and eigenvectors $\{\phi_r\}$ of a transition matrix **P** [1, 9]. Under an exchangeable graph, Lemma 2.1 proves that the diffusion maps at each t can provide conditional i.i.d samples $\{\mathbf{U}_t\}_{t\in\mathbb{N}}$.

Lemma 2.1 (Exchangeability and *i.i.d* of diffusion maps \mathbf{U}_t). Assume that \mathbf{G} is an exchangeable random graph that is connected, undirected and unweighted. Then the diffusion maps $\{\mathbf{U}_t(i): i=1,\ldots,n\}$ are conditionally *i.i.d* given its underlying distribution.

cs: are connected, undirected and unweighted graph really necessary? Yes, it should be connected and also undirected since coifman assumed symmetric adjacency matrix. Weighting usually does not matter but it matters in our case since proof of Lemme 2.1 is based on the assumption that A is binary.

Then we can define the *diffusion distance* between each pair of nodes, by computing the Euclidean distance of the diffusion maps.

$$C_t^2[i,j] := || \mathbf{U}_t(i) - \mathbf{U}_t(j) || \quad i,j = 1, 2, \dots, n.$$
 (1)

As diffusion time t increases, the corresponding diffusion distance C_t is more likely to take into account

of two nodes which are relatively difficult to reach each other. Figure 1 shows how diffusion distance can better reflects the connectivity and exhibits the community structure in a graph, when a reasonable t is chosen in the family of diffusion distance $\{C_t : t \in \mathbb{N}\}$. In practice, $t \in [5, 20]$ usually yields similar diffusion distance and suffices for later inferences. Compared to adjacent relation or geodesic distance which are two extremes in terms of the range of nodes in the count, diffusion distance well reflects the connectivity since it takes into account every possible path between the two nodes. cs: is the above sentence correct? Adjacent relation is equivalent to t = 0 and geodesic corresponds to $t = \infty$? : I did not mean adjacent relation necessarily corresponds to the case t = 0 and geodesic to $t = \infty$ but they are extreme in general.

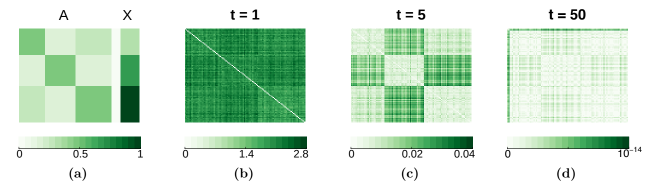


Figure 1: Figure (a) shows data generating probability of an adjacency matrix \mathbf{A} and nodal attributes \mathbf{X} . Diffusion matrix, as a proposed network metric, provides one-parameter family of network-based distances where as time goes by the pattern shown in the distance matrix changes, and at time point t = 5, distance matrix (c) illustrates most clear block structures and at the same time it exhibits most dependence to distance matrix of \mathbf{X} .

2.2 Dependence Testing via MGC

cs: why use W? why not just X or U(corresponding to diffusion map)? : I first started by introducing dCorr statistics so we did not have U. Here I will just leave it as (W,Y) since we already used up X as nodal attributes and we have subscript to U but not necessarily in general sample vector. The results in Section 2.1 allow us to cast the network dependency test into the following framework: given sample data $(\mathbf{U}, \mathbf{X}) = \{(\mathbf{u}_i, \mathbf{x}_i); i = 1, 2, ..., n\}$ that are identically distributed as $(\mathbf{u}, \mathbf{x}) \in \mathbb{R}^{D \times D_x}$ (D and D_x are the respective feature dimension), we are looking to test whether their joint distribution equals the product of the marginals, i.e.,

$$H_0: f_{\mathbf{u}\mathbf{x}} = f_{\mathbf{u}}f_{\mathbf{x}},$$

$$H_A: f_{\mathbf{1}\mathbf{1}\mathbf{x}} \neq f_{\mathbf{1}\mathbf{1}}f_{\mathbf{x}}.$$

If $(\mathbf{u}_i, \mathbf{x}_i)$ can be further assumed independently distributed for each i, a wide range of statistics are consistent for the above test, such as the distance correlation [19], HHG test [4], MGC statistic [16], etc. For example, the landmark distance correlation is computed as follows: denote $C_{ij} = ||\mathbf{u}_i - \mathbf{u}_j||$ and $D_{ij} = ||\mathbf{x}_i - \mathbf{x}_j||$ for i, j = 1, 2, ..., n, where $||\cdot||$ is the Euclidean distance. The distance covariance is defined as

$$dCov(\mathbf{U}, \mathbf{X}) = \frac{1}{n^2} \sum_{i,j=1}^{n} \tilde{C}_{ij} \tilde{D}_{ij}, \qquad (2)$$

where \tilde{C} and \tilde{D} is doubly-centered C and D by its column mean and row mean respectively, i.e., $\tilde{C} = HCH$, where $H = I_n - \frac{J_n}{n}$ (the double centering matrix), I_n is the $n \times n$ identity matrix (ones on the diagonal, zeros elsewhere), and J_n is the $n \times n$ matrix of all ones. The distance correlation dCorr follows by normalizing the distance covariance and is in the range of [0,1]. The best property of distance correlation is its consistency against almost all alternatives, i.e., dCorr(U, X) has testing power 1 for n large, for any joint distributions of finite moment. The MGC test inherits the consistency of distance correlation, and significantly improves the finite-sample testing power via locating the optimal local correlation, i.e., excluding far away distances in the computation of distance correlation.

However, as the i.i.d. assumption is not satisfied under network topology, the consistency of distance correlation is no longer guaranteed when applied to arbitrary distance metric of the graph. For example, neither the Euclidean distance of the adjacency vector nor the shortest-path distance can work together with distance correlation without breaking its consistency proof. Based on Lemma 2.1, we are able to prove that the both dCorr and MGC defined on the diffusion distance have the same consistency when extended to network dependency test.

Theorem 2.2. Assume that **G** is an exchangeable random graph that is connected, undirected and unweighted with n vertices with the diffusion maps **U** at certain t; and the nodal attributes $\mathbf{X} = \{\mathbf{x}_i, i = 1, 2, ..., n\}$ is i.i.d. as a random vector \mathbf{x} of finite moment.

Then $dCorr(\mathbf{U}, \mathbf{X}) \longrightarrow 0$ as $n \to \infty$ if and only if \mathbf{U} is independent of \mathbf{X} . Then both dCorr and MGC are consistent for testing dependence between \mathbf{U} and \mathbf{X} .

cs: not sure if thm 2 is necessary or not, excluded for now. : Yes, it might be redundant in this papar. I think I would include it in arxiv version since I applied MGC to FH network factors.

Therefore, we successfully extend distance-based correlation measures to the graph domain, offering a principal approach to define correlations and testing dependency on network data. We will next investigate our approach via simulated models and empirical performances.

2.3 Measure for Node Contribution

Can we just include MGC statistic? Without explaining (k,l) it becomes confusing to explain node contribution. On the other hand, some nodes often exert more reliance on their attributes than the others. Here we suggest the measure of node's contribution to detecting dependence as a byproduct of MGC statistic. Let (k^*, l^*) be the optimal neighborhood choice in distance matrix (C, D) respectively. Denote the contribution of node $v \in V(G)$ to the testing statistic by $c(\cdot) : v \to \mathbb{R}$

$$c(v) \propto \sum_{j=1}^{n} \tilde{C}_{jv} \tilde{D}_{jv} I\left(r(C_{jv}) \le k^*\right) I\left(r(D_{jv}) \le l^*\right), \tag{3}$$

which is proportional to v^{th} column-sum of the pre-summed test statistic of MGC. Note that the deviation of non-negative MGC statistic from zero implies departure from the independence and also note that we truncate the correlation in dCov by column entry's rank. Thus $\tilde{C}_{jv}\tilde{D}_{jv}$ would not be truncated if node j ($\in \{1, 2, ..., n\} \setminus \{v\}$) is important to node v and its larger, positive value would contribute to the statistic more. The statistic c(v) comes out from these observations.

3 Simulation Study

In our simulation studies, we compare the empirical testing powers of MGC, mCorr, Heller-Heller-Gorfine (HHG), and likelihood ratio test of Fosdick and Hoff (FH) [2]. For each simulation model and each statistic, we repeatedly generate sample data for 500 times, carry out the permutation test, and reject the null if the resulting p-value is less than 0.05. The testing power of each method equals the percentage of correct rejection.

3.1 Stochastic Block Model

As a common dependency structure in clustering or classification of vertices, in this simulation we assume the nodal attributes are distributed as \mathbf{x} , taking value in 0, 1, 2 equally likely. cs: I am confused on the model below, let us talk tmr We first consider a SBM with 3 blocks (equation 4) where block affiliation for each node is correlated with its attributes X. Assume that for $i, j = 1, \ldots, n = 100$, we have

$$E(A_{ij}|X_i, X_j) = 0.5I(|X_i - X_j| = 0) + 0.2I(|X_i - X_j| = 1) + 0.3I(|X_i - X_j| = 2).$$
(4)

these two nodes are most likely to have an edge but when X_i and X_j differ by one, they are even less likely to have an edge, with probability of 0.2, than the most different pairs of nodes. This actually describes nonlinear dependence where MGC is believed to work better than the distance correlation. Figure 2 illustrates that MGC combined with diffusion maps (DF) indeed yield the most superior power than others.

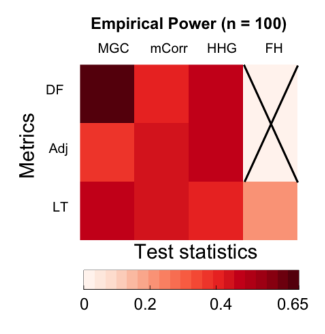


Figure 2: This power heatmap illustrates the superior power of multiscale generalized correlation (MGC) under diffusion distance matrix (DF) in three-block SBM (equation 4), compared to under adjacency matrix distance (Adj) or latent factor distance (LT). This demonstrates one exemplary network where MGC statistic along with a family of diffusion distances catches non monotonic correlations efficiently than the other statistics and metrics.

To scrutinize our conjecture on better performance of local optimal scaled MGC over global scale of mCorr, we control the amount of nonlinear dependency through changing the value of $\theta \in (0,1)$ in the three block equation 5.

 $E(A_{ij}|X_i,X_j)=0.5I(|X_i-X_j|=0)+0.2I(|X_i-X_j|=1)+\theta I(|X_i-X_j|=2), \quad i,j=1,\ldots,n=100, \ (5)$ where $X_i \overset{i.i.d}{\sim} Bern(0.5); i=1,\ldots,100.$ When $\theta>0.2$, linear dependency of edge distribution in $\mathbf A$ upon nodal attribute of X is lost. If you see Figure 3, power of mCorr starts to drop from $\theta=0.2$ while that of MGC almost stays clam, which implies MGC is significantly more sensitive to nonlinear dependency compared to mCorr.

On the other hand, the SBM connotes that all nodes within the same block have the same expected degree. Thus, this block model is limited by homogeneous distribution within block and provides a poor fit to networks with highly varying node degrees within block or community. Instead the Degree-Corrected Stochastic Block model (DCSBM) proposed by [8] add another random variable associated with each node to vary the node degrees. In the model 6, we controlled the amount of such variability by τ ; the larger the value τ is, the more variability degree or edge distribution has. In Figure 4, power

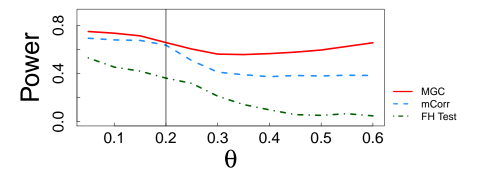


Figure 3: X-axis of θ controls the existence/amount of nonlinear dependency and in this particular case nonlinearity exists when $\theta > 0.2$ and gets larger as it increases. You can see the discrepancy in power between global and local scale tests also gets larger accordingly, mostly due to decreasing power of mCorr or FH test but relatively stable power of MGC under nonlinear dependency.

based on Euclidean distance of $\bf A$ or that of estimated network factors (locations) becomes less sensitive as τ increases. Compared to these two, diffusion maps are more robust to such variability.

$$E(A_{ij}|\mathbf{X}, \mathbf{V}) = 0.2V_i V_j \cdot I(|X_i - X_j| = 0) + 0.05V_i V_j \cdot I(|X_i - X_j| = 1), \tag{6}$$

where $X_i \stackrel{i.i.d.}{\sim} Bern(0.5); V_i \stackrel{i.i.d.}{\sim} Uniform(1-\tau, 1+\tau), i=1,\dots, 250.$

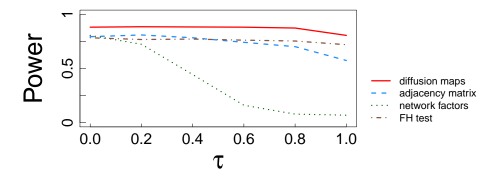


Figure 4: In degree-corrected SBM where the variability in degree distribution increases as τ increases, testing power of diffusion maps are more likely to be robust against increasing variability compared to other network metrics, e.g. adjacency matrix or latent positions. FH test statistics allowing different dimensions of network factors perform consistently well but still have less power than MGC.

3.2 Node Contribution Test

To examine the effectiveness of node contribution measure in testing dependency as presented in the statistic 3, we deliberately simulate the network and its nodal attributes as half of the nodes are

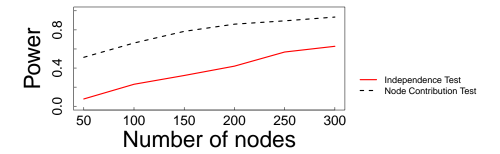


Figure 5: This plot describes that both power of MGC and the rate of correctly-ranked node contribution increase as the number of nodes increases when only half of the nodes for each simulation actually are set to be dependent on network, which validates the use of node contribution measure in independence test.

independent while the other half are dependent on network (model 7).

$$X_{i} \stackrel{i.i.d}{\sim} Bern(0.5) \quad i = 1, \dots, n/2, \dots, n$$

$$E(A_{ij}|X_{i}, X_{j}) \stackrel{d}{=} \begin{cases} 0.4I(|X_{i} - X_{j}| = 0) + Bern(0.1)I(|X_{i} - X_{j}| > 0) & i = 1, \dots, n/2 \\ 0.25 & i = 1 + n/2, \dots, n \end{cases}$$

$$(7)$$

As an ad hoc test of node contribution, we rank the nodes in terms of decreasing order of c(v) and count the ratio of dependent samples's ranks within the number of dependent nodes. If it works perfectly, all dependent nodes would take higher rank than every independent node so thus the rate equals to one. We call this rate as *inclusion rate*:

inclustion rate
$$(c(v)) = \sum_{v \in V(\mathbf{G})} \{rank_{c(v)}(v) \le m\}/m,$$
 (8)

where $m(\leq |V(\mathbf{G})|)$ is the number of nodes under network dependence. We set m = n/2 out of $n = |V(\mathbf{G})|$.

4 Conclusion

In this paper, we convince that MGC, merged with a family of diffusion distance, provides us powerful independence test statistics in network. Having multiscale statistics, i.e. one parameter family of statistics, is not avoidable because we regard distance between the nodes over network as a dynamic process. Through simulation studies, we demonstrate that our methods perform better than the others especially under nonlinear dependency, and we are able to measure each node's contribution to detecting dependency. Deriving the contributions is particularly important when there have possibly different amounts

of the dependencies among the nodes.

However obtaining a full family of statistics are computationally infeasible. Also we did not suggest any theoretically supported tools to select one metrics among them so thus we have one single statistic. As an ad hoc, we selected an *optimal* diffusion time t with highest power from t=1 to t=10 for our simulation since we could observe a stabilized empirical power within this period. Developing the adaptive method to find this optimal t where dependence is maximized would be a natural next step. Despite these shortcomings, we expect that we could also enjoy the properties of MGC and a family of diffusion distances in solving diverse problems which require to utilize local relationship of the data sets. For instance, we might be able to implement independence testing between two networks of same size by using diffusion distance of each network to investigate whether a pair of networks are topologically or structurally independent. This kind of work would shed light on revealing any relationship between the data sets which are not necessarily a random vector.

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5 Appendix

5.1 Lemmas and Theorems

Proof of Lemma 2.1. Diffusion map at time t is represented as follows:

$$\mathbf{U}_t(i) = \begin{pmatrix} \lambda_1^t \phi_1(i) & \lambda_2^t \phi_2(i) & \cdots & \lambda_q^t \phi_q(i) \end{pmatrix} \in \mathbb{R}^q.$$
 (9)

where $\Phi = \Pi^{-1/2}\Psi$ and $Q = \Psi\Lambda\Psi^T = \Pi^{1/2}P\Pi^{-1/2}$. Thus $P\Pi^{-1/2}\Psi = \Pi^{-1/2}\Psi\Lambda$. Then for any rth row $(r \in \{1, 2, ..., q\}, (q \leq n))$, we can see that $P\phi_r = \lambda_r\phi_r$ where $\phi_r = \left(\psi_r(1)/\sqrt{\pi(1)} \ \psi_r(2)/\sqrt{\pi(2)} \ \cdots \ \psi_r(n)/\sqrt{\pi(n)}\right)$. Therefore to guarantee exchangeability (or i.i.d) of \mathbf{U}_t , it suffices to show exchangeability (or i.i.d) of P.

Assume joint exchangeability of **G**, i.e. $(A_{ij}) \stackrel{d}{=} (A_{\sigma(i)\sigma(j)})$. Since A_{ij} is binary, $A_{ij}/\sum_j A_{ij} = A_{ij}/(1+\sum_{l\neq j}A_{il})$. Moreover, A_{ij} and $(1+\sum_{l\neq j}A_{il})$ are independent given its link function g, and $A_{\sigma(i)\sigma(j)}$ and $(1+\sum_{l\neq j}A_{\sigma(i)\sigma(l)})$ are independent also given g. Then the following joint exchangeability of transition probability holds for $i\neq j; i,j=1,2,\ldots,n$:

$$\left(P_{ij}\right) = \left(\frac{A_{ij}}{1 - A_{ij} + \sum_{j=1}^{n} A_{ij}}\right) \stackrel{d}{=} \left(\frac{A_{\sigma(i)\sigma(j)}}{1 - A_{\sigma(i)\sigma(j)} + \sum_{\sigma(j)=1}^{n} A_{\sigma(i)\sigma(j)}}\right) = \left(P_{\sigma(i)\sigma(j)}\right) \tag{10}$$

When i = j, $P_{ij} = P_{\sigma(i)\sigma(j)} = 0$ for i = 1, 2, ..., n. Thus, transition probability is also exchangeable. This results exchangeable eigenfunctions $\{\Phi(1), \Phi(2), ..., \Phi(n)\}$ where $\Phi(i) := \begin{pmatrix} \phi_1(i) & \phi_2(i) & \cdots & \phi_q(i) \end{pmatrix}^T$,

 $i=1,2,\ldots,n$. Thus diffusion maps at fixed t, $\mathbf{U}_t=\begin{pmatrix} \Lambda^t\Phi(1) & \Lambda^t\Phi(2) & \cdots & \Lambda^t\Phi(n) \end{pmatrix}$ are exchangeable. Furthermore by $de\ Finetti's\ Theorem$, we can say that $\mathbf{U}(t)=\{\mathbf{U}_t(1),\mathbf{U}_t(2),\ldots,\mathbf{U}_t(n)\}$ are conditionally independent on their underlying distribution.

Proof of Theorem 2.2 Consistency of dCorr applied to exchangeable variables. For exchangeable sequence of $(\mathbf{U}, \mathbf{X}) = \{(\mathbf{u}_i, \mathbf{x}_i); i = 1, 2, ..., n\}$ which is identically distributed as (\mathbf{u}, \mathbf{x}) with finite second moment, we have

$$\mathcal{V}_n^2(\mathbf{U}, \mathbf{X}) \longrightarrow \mathcal{V}^2(\mathbf{u}, \mathbf{x}) \quad \text{as } n \to \infty$$
 (11)

where $\mathcal{V}^2(\mathbf{u}, \mathbf{x}) := \|g_{\mathbf{u}, \mathbf{x}}(t, s) - g_{\mathbf{u}}(t)g_{\mathbf{x}}(s)\|^2$, and g is a characteristic function, e.g., $g_{\mathbf{u}, \mathbf{x}}(t, s) = E\{\exp\{i\langle t, \mathbf{u}\rangle + i\langle s, \mathbf{x}\rangle\}\}$. This follows exactly the same as *Theorem 1* in [19]. Note that this Lemma always holds without any assumption on $\{(\mathbf{u}_i, \mathbf{x}_i), i = 1, 2, ..., n\}$.

Followed by de Finetti's Theorem, if and only if $\{\mathbf{u}_i\}$ are (infinitely) exchangeable, there exists an underlying distribution $f_{\mathbf{u}}$ of \mathbf{u} such that $\mathbf{u}_i \stackrel{i.i.d}{\sim} f_{\mathbf{u}}$. By the same logic there exists a random, we have an underlying distribution $f_{\mathbf{x}}$ where $\mathbf{x}_i \stackrel{i.i.d}{\sim} f_{\mathbf{x}}$. Let $(\mathbf{u}_i, \mathbf{x}_i) \stackrel{i.i.d}{\sim} f_{\mathbf{u},\mathbf{x}}$. Then under the assumption of finite second moment of the underlying distributions and measurable, conditioned random functions, we have a strong large number for V-statistics followed by [19], i.e.,

$$\int_{D(\delta)} \|g_{\mathbf{u},\mathbf{x}}^n(t,s) - g_{\mathbf{u}}^n(t)g_{\mathbf{x}}^n(s)\|^2 dh \stackrel{n \to \infty}{\longrightarrow} \int_{D(\delta)} \|g_{\mathbf{u},\mathbf{x}}(t,s) - g_{\mathbf{u}}(t)g_{\mathbf{x}}(s)\|^2 dh, \tag{12}$$

where $D(\delta) = \{(t,s) : \delta \le |t|_p \le 1/\delta, \delta \le |s|_q \le 1/\delta\}$, and h(t,s) is the weight function chosen in [19]. It follows that

$$\mathcal{V}_n^2(\mathbf{U}, \mathbf{X}) \to 0 \quad \text{as } n \to \infty$$
 (13)

if and only if $g_{\mathbf{u},\mathbf{x}}(t,s) = g_{\mathbf{u}}(t)g_{\mathbf{x}}(s)$, i.e., \mathbf{u} is independent of \mathbf{x} . Therefore, the dCorr or mCorr converges to 0 if and only if underlying distributions are independent; and its testing power converges to 1 under any joint distribution of finite moments. Since the multiscale generalized correlation based on any consistent global correlation is also consistent [16], MGC statistic constructed by dCorr or mCorr is also consistent in testing dependence.