# Statistical Inference of Discrete Combinatorial Functional Dependency in Biological Systems 

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#### Abstract

Inference of a combinatorial function from multiple independent variables (parents) to a dependent variable (child) in a discrete space can be useful in detecting nonlinear relationships in biological systems. Popular conditional independency measures, heavily used in combinatorial inference, are often insensitive to the direction of functional dependency. To address this issue, we define multivariate and conditional functional chi-squared statistics. We also present an algorithm called CFDF for bivariate discrete function inference via an exclusive-effect strategy, in order to identify a best parent set for a given child. It requires each parent to make sufficient contribution beyond any marginal effect. Simulation studies suggest a marked advantage of our framework over alternatives. Applying the method to transcriptome data in genetically perturbed biological systems, we reproduced combinatorial gene interactions known in the literature. Most importantly, we identified combinatorial patterns from joint RNA and protein data to rebut a dispute on the founding principle of molecular biology.


## 1 Introduction

A combinatorial effect refers to the control of dependent (child) variable by multiple independent (parent) variables. Combinatorial control occurs widely in gene regulation [1, 2]; it drives exponential biodiversity with only linear increase in genetic materials [3]. A fruit fly survives only if it is either male with an active $S x l$ gene or female with an inactive $S_{x l}$ gene; in this example, survival is an exclusive-OR (XOR) function of sex and $S x l$ activity [4].
Many methods have been proposed to infer discrete combinatorial relationships, including switch-like models [5], Boolean networks [6, 7], discrete Bayesian networks [8] and their dynamic version [9], graphical models [10], Petri nets [11], and generalized logical networks [12]. However, these approaches often select one factor at a time [13, 14] and resort to linear [15] or sigmoidal [16] combinatorial functions, to gain computational efficiency at the cost of combinatorial interactions such as exclusive-OR. On the other hand, the more flexible conditional mutual information [17] can tell the difference between $X \not \perp Y \mid Z$ and $X \perp Y \mid Z$, and is used in inferring combinatorial transcription regulation [18]; however, it cannot infer $X \rightarrow Y \mid Z$ versus $X \perp Y \mid Z$, due to its symmetry between $X$ and $Y$.
Based on the top performing univariate functional chi-squared (FUNCHISQ) method [19] in causal biological network inference [20], we define multivariate FUNCHISQ for joint effects and conditional FUNCHISQ for combinatorial effects. The framework is based on the exclusive-effect premises: 1) the combinatorial effect is captured by conditional statistics exclusive of the marginal effect and 2) when no marginal effect in a given data set can fully explain a child variable, this data set may afford a chance to capture a combinatorial effect. We give a branch-and-bound algorithm called CFDF to
identify a best bivariate discrete function and demonstrate its effectiveness on both simulated and biological data sets.

## 2 Methods

Definition 1 (Univariate FUNCHISQ statistic). Given $n$ samples of discrete random variables $X$ and $Y$ of $q$ and $Q$ levels, respectively, the univariate FUNCHISQ is [19, 21]

$$
\begin{equation*}
\chi_{f}^{2}(X \rightarrow Y)=\left[\sum_{i=1}^{q} \sum_{j=1}^{Q} \frac{\left[n_{i j}-\left(n_{i \cdot} / Q\right)\right]^{2}}{n_{i \cdot} / Q}\right]-\left[\sum_{j=1}^{Q} \frac{\left[n_{\cdot j}-(n / Q)\right]^{2}}{n / Q}\right] \tag{1}
\end{equation*}
$$

where $n_{i j}$ is the count of $X=i$ and $Y=j$ in the $q \times Q$ contingency table whose rows represent $X$ and columns $Y, n_{i}$. is the sum of row $i$, and $n_{\cdot j}$ is the sum of column $j$.

It was previously established that $\chi_{f}^{2}(X \rightarrow Y)$ asymptotically follows a chi-square distribution of $(q-1)(Q-1)$ degrees of freedom under the null hypothesis that $Y$ is independent of $X$ and the assumption that $Y$ is uniformly distributed [19]. A related exact functional test based on the same test statistic is also established [22]. FUNCHISQ is unique in asymmetric functional optimality and is maximized if and only if $Y$ is a non-constant function of $X$ given $Y$ 's marginal [23]. The statistic is minimized to zero when $X$ and $Y$ are empirically independent. FUNCHISQ previously outperformed mainstream techniques for causal biological network inference [20].
Definition 2 (Multivariate FUNCHISQ statistic). Let $X=\left[X_{1}, \ldots, X_{p}\right]^{\top}$ be a p-dimension discrete random vector with $q_{1}, \ldots, q_{p}$ levels, respectively. Let $Y$ be a discrete random variable of $Q$ levels. We construct a contingency table of $q=q_{1} \times \cdots \times q_{p}$ rows and $Q$ columns. For an observed vector of $x=\left[x_{1}, \ldots, x_{p}\right]^{\top}$, we linearize it to calculate a scalar row index $i$ by

$$
\begin{equation*}
i=\left[\sum_{k=1}^{p-1}\left(x_{k} \prod_{l=k+1}^{p} q_{l}\right)\right]+x_{p} \tag{2}
\end{equation*}
$$

Plugging $q$ and i into Eq $\sqrt{17}$, we define the multivariate FUNCHISQ statistic $\chi_{f}^{2}\left(X_{1}, \ldots, X_{p} \rightarrow Y\right)$, which is the joint effect of $X$ on $Y$.

The multivariate FUNCHISQ is also asymptotically chi-squared distributed under the null hypothesis that $Y$ is independent of $X$ and the assumption that $Y$ is uniformly distributed, following a similar argument in [19]. The statistic is subject to spurious combinatorial relationship, as it may pick a parent set involving only one strong parent. To overcome this issue, we introduce conditional functional chi-squared statistic to determine sensible combinations of parents in a multivariate functional relationship.
Definition 3 (Conditional FunChisQ statistic). Let $X=\left[X_{r}, X_{s}\right]^{\top}$. We define the conditional functional chi-squared statistic from random variable $X_{s}$ to $Y$ given $X_{r}$ by

$$
\begin{equation*}
\underbrace{\chi_{f}^{2}\left(X_{s} \rightarrow Y \mid X_{r}\right)}_{\text {combinatorial effect }}=\underbrace{\chi_{f}^{2}(X \rightarrow Y)}_{\text {joint effect }}-\underbrace{\chi_{f}^{2}\left(X_{r} \rightarrow Y\right)}_{\text {marginal effect }} \tag{3}
\end{equation*}
$$

It can be shown that $\chi_{f}^{2}\left(X_{s} \rightarrow Y \mid X_{r}\right)$ follows an asymptotic chi-squared distribution with $q_{r}\left(q_{s}-\right.$ 1) $(Q-1)$ degrees of freedom under the null hypothesis that $Y$ is independent of $X_{r}$ and $X_{s}$ and the assumption that $Y$ is uniformly distributed. Conditional FUNCHISQ promotes combinatorial patterns with a large difference between joint and marginal effects. It promotes parents that perform poorly by themselves but outperform best individual parents when combined. For example, in an XOR relationship, looking at $X_{1}$ with respect to $Y=X_{1} \oplus X_{2}$ results in a poor individual (marginal) effect; however, looking at $X_{1}, X_{2}$ together maximizes the joint effect.
The combinatorial functional dependency by FUNCHISQ (CFDF) algorithm (Algorithm 1). It selects a best bivariate combinatorial relationship $X_{i}, X_{j} \rightarrow Y$ among pairs in $X$ of $v$ variables. It uses a branch-and-bound strategy similar to the $A^{*}$ search. It maintains $\beta \in[0,1]$, the lowest combinatorial (conditional FUNCHISQ) $p$-value so far and $\chi_{\max }^{2}$, the maximum joint FUNCHISQ

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Algorithm 1 CFDF: Combinatorial Functional Dependency by FUNCHISQ
Input: Child \(Y\) and parent candidates \(\mathbf{X}=\left[X_{1}, X_{2}, \ldots, X_{v}\right]\)
Output: A best pair of parents \(P_{1}, P_{2} \in \mathbf{X}\)
    Calculate univariate FunChISQ statistics (individual effects) of each \(X_{i}\) in \(\mathbf{X}\)
    Sort \(\mathbf{X}\) to \(\mathbf{X}^{\prime}\) by individual effect in non-descending order
    Initialize \(\beta \leftarrow 1\) for minimum conditional FUNCHISQ \(p\)-value so far; \(\chi_{\max }^{2} \leftarrow 0\) for maximum joint FUNCHISQ achieved so far
    for each \(X^{(i)}\) in \(\mathbf{X}^{\prime}\) :
        for each \(X^{(j)}\) in \(\mathbf{X}^{\prime}\) and \(j>i\)
            Calculate optimistic conditional FUNCHISQ for both \(X^{(i)}, X^{(j)}\)
            if optimistic conditional FUNCHISQ \(p\)-values for both \(X^{(i)}, X^{(j)} \leq \beta\)
                \(p[i], p[j] \leftarrow\) actual conditional FunCHISQ \(p\)-values of \(\chi_{f}^{2}\left(X^{(i)} \rightarrow Y \mid X^{(j)}\right)\) and \(\chi_{f}^{2}\left(X^{(j)} \rightarrow Y \mid X^{(i)}\right)\)
            \(\beta \leftarrow \min (\beta, \max (p[i], p[j]))\)
            if \(\chi_{\max }^{2}<\chi_{f}^{2}\left(X^{(i)}, X^{(j)} \rightarrow Y\right) \#\) collective effect of \(X^{(i)}, X^{(j)}\)
                \(\chi_{\text {max }}^{2} \leftarrow \chi_{f}^{2}\left(X^{(i)}, X^{(j)} \rightarrow Y\right)\)
                \(P_{1} \leftarrow X^{(i)}\) and \(P_{2} \leftarrow X^{(j)}\)
            else if optimistic conditional FUNCHISQ of \(X^{(j)}>\beta\)
            Remove \(\left[X^{(j)} X^{(j+1)}, \cdots, X^{(v)}\right]\) from \(\mathbf{X}^{\prime}\)
            Terminate the inner for-loop
        else if optimistic conditional FUNCHISQ of \(X^{(i)}>\beta\)
            Terminate the outer for-loop
    Return \(P_{1}, P_{2}\)
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statistic so far. The optimistic conditional FUNCHISQ is the $p$-value associated with the theoretical maximum of the statistic, given the individual effect. The returned parents of $Y$ have a large joint effect constrained by sufficient conditional effects. The algorithm has a worst-case runtime of $O\left(v^{2}\right)$. Speedup occurs when a variable fails to secure an optimistic contribution better than or equal to $\beta$, in which case we terminate the outer loop if the variable was $X^{(i)}$, since any other $X^{(k)}, k>i$ in the ordered list cannot do better; and in case of $X^{(j)}$, we remove all $X^{(k)}, k \geq j$ from the ordered list followed by inner loop termination.

## 3 Results

Simulation studies. To evaluate the performance of CFDF, we randomly generated 10 independent variables and constructed 20 bivariate functions from polynomial, trigonometric, logarithmic and exponential function families. All 10 parents and 20 children were discretized using optimal univariate clustering from the 'Ckmeans.1d.dp' package [24, 25], with the number of clusters determined using the 'mclust' package [26]. We generated 1500 samples with noise added by the R function jitter.
We compared CFDF to four other discrete conditional independency tests from the 'bnlearn' package [27] including conditional mutual information, conditional Pearson's chi-square, semi-parametric conditional mutual information and semi-parametric conditional Pearson's chi-square. Each method evaluated all 30 variables as a potential child variable to return a ranked list of the remaining 29 variables, ideally ranking the true parents (when present) higher than others, distinguishing a functional from an independent relationship and also predicting the parent $\rightarrow$ child direction. Figure $1 \mathrm{a}, \mathrm{b}$ summarize AUPR and AUROC over multiple runs of the same setup with increasing noise. CFDF performs similarly in AUROC but outperforms other strategies in AUPR, suggesting that CFDF better ranks true directional interactions, as CFDF was the only directional measure out of all competitors.

Pattern discovery on leukemia gene expression data. We applied CFDF on two real data sets for both validation and discovery of combinatorial interactions. All molecular quantities were log transformed and discretized using 'Ckmeans.1d.dp' [24], with quantization levels determined by 'mclust' package [26]. We used the immortalized K562 leukemia cancer cell-line RNA-seq data collected after the knockdown of $\sim 250$ RNA-binding proteins (RBPs) from the ENCODE project [28]. Each sample measured the abundance of $\sim 195 \mathrm{~K}$ transcripts in transcripts per million. For each RBP, we searched for its best combinatorial parents amongst the remaining RBPs. Figure 1 c shows a known interaction that we were able to validate from literature [29-31]. More encouragingly, Figure 1d shows a putative combinatorial interaction close to an exclusive-OR function.
Rebutting a dispute on the central dogma of molecular biology. The central dogma of molecular biology states that "DNA makes mRNA and mRNA makes protein" [32]. While a good correlation has been observed between DNA copy number and mRNA expression of the same gene [33-35], recent joint mRNA and protein studies [33-35], reported low correlation between the mRNA and protein levels of a gene, challenging the central dogma. We offer alternative evidence to show that the central dogma is not violated and the observed weak mRNA-protein correlation can be explained by


Figure 1: Result summary. (a) AUPR and (b) AUROC over multiple runs with increasing jitter noise. (c) PHF6 $\times$ CNOT7 $\rightarrow$ NPM1 and (d) CCAR $1 \times$ DDX24 $\rightarrow$ DAZAP1, where the X-axis is the first parent, Y-axis the child, and color represents the second parent. The dotted lines represent the quantization levels. (e) and (f) C2orf49, TCEA1's mRNA and protein form a strong combinatorial relationship ( $P=2.6 \times 10^{-5}$ ), rebutting central dogma violation.
combinatorial gene regulation. The Clinical Proteomics Tumor Analysis Consortium [35] provided 3718 proteins of 30 matched normal and 90 colorectal tumor samples from The Cancer Genome Atlas (TCGA) project. About 57000 mRNA transcripts for matched samples were extracted from TCGA. For each of the 3718 proteins, univariate FUNCHISQ between a protein and its mRNA was computed. If the mRNA was found to be a bad predictor for the protein ( $p$-value $>0.05$ ), this protein was taken to the CFDF algorithm with one parent fixed to be the mRNA of the protein. CFDF then searched for a possible second mRNA/protein for a combinatorial effect.
We found a strong combinatorial pattern in translation of gene TCEA1 mRNA to its protein involving a second gene C2orf49 (Figure 11). The mRNA and protein of TCEA 1 are only weakly correlated (Figure 1e). However, a strong pattern emerges if we consider C2orf49's mRNA in three levels: at each level, a curve exhibits different translation rates and basal levels for TCEA1. Thus, given the mRNA level of C2orf49, the mRNA level of TCEA1 indeed predicts the protein level of TCEA1. TCEA1 and C2orf49 are related as belonging to the gene expression SuperPath from PathCards [36]. As translation may involve multiple factors, our finding supports that combinatorial effects reinforce rather than violate the central dogma of molecular biology.

## 4 Conclusions

The FUNCHISQ based combinatorial functional inference can capture both nonlinear and nonmonotonic functional relationships. CFDF selects sensible combinations of variables by requiring sufficient contributions from each parent variable to the child variable. Our simulation studies demonstrate marked improvement of CFDF over alternative methods on a variety of noisy multivariate functions. We uncovered combinatorial patterns to rebut a dispute on a basic principle of molecular biology. As the method does not assume a predefined functional model, it is applicable to combinatorial pattern discovery in many under-studied biological systems.

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