Multi-task Recurrent Neural Network and Higher-order Markov Random Fields for Stock Price Movement Direction Prediction

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ABSTRACT

Stock price movement not only depends on individual stocks' historical records, but also has complex hidden dynamics associated with other correlated stocks. Despite a substantial amount of effort has been made to understand the principles of stock price movement, few has attempted to predict movement direction based upon single stock's historical records as well as its correlated stocks. To this end, we present a multi-task recurrent neural network (RNN) and high-order Markov random fields (MRFs) to perform stock price movement direction prediction. Specifically, we first design a multitask RNN framework to extract informative features from the raw market data of individual stock without considering any domain knowledge. Next, we employ binary MRFs with unary as well as weighted lower linear envelope as the higher-order energy function to capture the higher-order consistency within the same clique (group) of stocks. We also derived a latent structural SVM algorithm to learn high-order MRFs in a polynomial number of iterations. Finally, a sub-gradient algorithm is employed to perform end-to-end training of RNN and high-order MRFs. We conduct thoroughly empirical studies based on three popular Chinese stock market indexes and the proposed method outperforms baseline approaches. To the best of our knowledge, the proposed technique is the first one to investigate intra-clique relationships with higher-order MRFs on stock price prediction.

CCS CONCEPTS

• Computer systems organization \rightarrow Embedded systems; *Redundancy*; Robotics; • Networks \rightarrow Network reliability.

KEYWORDS

Deep learning, Graphical model, Multivariate time series prediction

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1 INTRODUCTION

It has been well-known that single stock's price movement not only depends on its historical records, but also is highly correlated to other stocks [27, 30] and may change in a non-synchronous manner [6, 27]. This correlated yet asynchronous price movement phenomenon is sometimes defined as lead-lag relationship [17] within a group of stocks. Different speed of information diffusion has been believed to be the main reason of lead-lag relationship [2, 27, 29]. When an information hits the market, some stocks'price tend to react faster than others. Therefore, identification of those leading stocks and their lead-lag relationships to other lagging stocks will provide strong evidence in predicting the latter's price movement direction.

There are three key challenges in utilizing the lead-lag relationship: (1) Discovering which stock will be affected by newly arrived information; (2) Identifying the group (e.g., industry, supply chain, etc.) it belongs to as well as leading stocks and lagging stocks in this group. Then modeling their relationships; (3) Predicting price movement of each stock by jointly considering the knowledge in its correlated group and individual stock's price movement at current stage.

The first challenge is extremely difficult to resolve with an automatic algorithm. This is not only because it requires an expert level understanding of the finance system and the dynamics behind market news as well as stock price, but also due to lacking of training data. However, according to the well-known efficient market hypothesis [28], stock price reflects all available market information. Therefore, informative stock price changes can be employed as an proximity to market news arrival. In this way, the complexity of the first challenge is transformed to the detection of informative price changes in individual stock.

Economists have spent decades trying to use patterns hidden inside historical trading price and volume to predict future price movement [10, 20]. Those models are called technical analysis [21]. However, most of those models have been proven stopped generating profitable signals since the early 1990s [32]. On the other hand, since trading strategies based on technical analysis rules are public available and easy to replicate, informed institutional traders are motivated to manipulate market price and trap retail (individual) traders following their manipulated price to gain excess profit [37].

To overcome those problems and address the first challenge, we employ an end-to-end hierarchical multi-task [7] RNN to extract informative changes from raw market price without using any of hand-crafted features such as technical analysis indicators. Good performance on price prediction relies on rich representations as

well as the multi-task framework which can leverage complementary aspects from diverse tasks [36]. Specifically, given the raw market price data which only contains six features (*i.e.*, opening price, low price, high price, closing price, volume, and amount) at each time interval, we leverage a hierarchical multi-task network to extract features on different tasks first and then concatenate those complementary feature vectors to make final prediction.

To model lead-lag relationships and address the other two challenges, we present a binary Markov Random Fields (MRFs) with weighted lower linear envelope as higher order (when the clique contains more than two nodes) energy function [12, 13, 23, 31]. In our implementation, we treat each stock as a node in MRFs and each stocks group which has lead-lag relationships as a maximum clique in MRFs. We use pre-defined industry classification list [1] as the prior domain knowledge of each stock's belonging maximum clique. Because we use a weighted version of higher order function, stocks have higher weights in the above list can be seen as leading stocks, and vice versa. Finally, the complexity of modeling dynamics between leading and lagging stocks becomes encouraging consistent over large cliques under weighted lower linear envelope potentials. Logits from hierarchical RNN networks are used as unary features in MRFs. By minimizing the energy function which contains both unary and higher order features, we are able to predict each stock's future price movement by jointly considering individual market price trending together with lead-lag relationships.

Unlike the first challenge trying to avoid prior knowledge, here we consider being able to embed prior knowledge as an advantage of our model. Definitions of sectors as well as leading and lagging stocks in each sector require solid financial industry research. Statistical evidences learned automatically from market price data are usually insufficient for determining such relationships.

We justify the effectiveness of the proposed technique based on three popular Chinese stock market indexes and the proposed method outperforms baseline approaches. To the best of our knowledge, the proposed technique is the first one to investigate intraclique relationships with higher-order MRFs on stock price prediction.

To summarize, the main contributions of this paper include:

- We propose a hierarchical multi-task RNN architecture to learn stock price patterns without any hand-crafted feature.
- We propose the first model that encode lead-lag relationship among stocks using higher-order MRFs.
- We develop an algorithm to learn weighted lower linear envelope with latent variables as higher order energy function under latent structural SVMs framework. Adding latent variable into higher order function enables our model to learn much richer representation than previous study [13].

2 RELATED WORKS

This work is closely related to lead-lag relationship, multi-task learning, high-order MRFs, as well as latent structural SVMs.

Lead-lag relationship Lead-lag relationship has been found to be a long existence phenomenon in stock market. Many reasons can lead to it such as diffusion of information, sector (industry) rotation, investment style rotation, event driven trading and asynchronous trading [8, 9, 14, 27]. Generally it is believed that lead-lag

relationship is more prevalent for firms in the same industry [17]. This assumption gives rise to our setting that we use pre-defined industry classification list [1] as prior domain knowledge of each stock's belonging maximum clique. A lot of research [2, 6, 17, 29] indicates that stocks with larger capital size and higher liquidity tend to be leading stocks and vice versa. To replicate potential lead-lag relationship, we assign each stock a different weight from its corresponding indexes created by the China Securities Index Company, Ltd. More complicated dynamics hidden behind clique of stocks are learned by higher-order MRFs.

Multi-task learning Caruana [7] showed that inductive knowledge learned from multiple tasks can transfer between tasks and help improving generalization on all tasks. Many works in Natural Language Processing (NLP) area take advantage of multi-task framework and achieves state-of-the-art performance while using simple models for each of these tasks [16, 36]. However, as pointed out by many researchers [7, 35], there are lacking of theories on choosing a diverse set of tasks and hierarchical architecture of chosen tasks. Recent works [16, 36] apply the intuition that the complexity of task should be increasing along with hierarchical level. We follow this intuition in our implementation. Because technical analysis indicators can be categorized into four categories (trend, momentum, volatility and volume) [21] and volume is included in market price data, we propose an architecture that using trend and volatility tasks as our lower level tasks and price movement classification (upward or downward) as higher level task. Other selection of tasks and hierarchical designations remain open for further research.

Higher-order Markov random fields Markov random fields are also known as undirected graphical model that can be seen as a regularized joint log-probability distribution of arbitrary nonnegative functions over a set of maximal cliques of the graph [3]. Utilizing MRFs mainly involves three steps: defining energy functions, solving inference problem (MAP or energy minimization) and learning parameters. As for energy functions, our work focus on a class of higher-order potentials defined as a concave piecewise linear function which is known as lower linear envelope potentials over a clique of binary variables. It has been raising much interest due to its capability of encoding consistent constraints over large subsets of pixels in an image [23, 31]. We follow Gould [13] to construct a graph-cut algorithm to solve exact inference problem and propose our novel learning algorithms under latent structural SVM in section 4.1.

As the second step, in order to solve inference problem, Kohli et al. [25] proposed a method to represent a class of higher order potentials with lower (upper) linear envelope potentials. By introducing auxiliary variables [22], they reduced the linear representation to a pairwise form and proposed an approximate algorithm with standard linear programming methods. However, they only show an exact inference algorithm on at most three terms. Following their routine, Gould [13] extended their method to a weighted lower linear envelope with arbitrary many terms which can be solved with an efficient algorithm. They showed the energy function with auxiliary variables is submodular by transforming it into a quadratic pseudo-Boolean form [4] and how graph-cuts [5, 11, 15] like algorithm can be applied to do exact inference.

As the third step, Gould [13] solved learning problem of lower linear envelope under the max margin framework [40]. In their work,

they pointed out the potential relationship between their auxiliary representation and latent SVM [43]. Our work is closely based on their research. We continue to use the higher order energy function and inference algorithm developed in their previous work [12] and extend their max margin learning algorithm to include latent variables. The learning algorithm we use is an extension of max margin framework which is known as "latent structural SVM" [43].

Latent structural SVMs The max-margin framework [39, 40] is a principled approach to learn weights of pairwise MRFs. Szummer et al. [38] adapted this framework to optimize parameters of pairwise MRFs inferred by graph-cuts method. In order to adapt higher-order energy function to max margin framework, Gould [12] approximated the energy function using equally spaced breakpoints. Gould [13] extended this framework with additional linear constraints which enforces concavity on weights thus can be used for learning lower linear envelope potentials. However, those methods can only learn higher-order functions approximately. In this paper we propose an algorithm to optimize the energy function exactly by introducing auxiliary variables back into the feature vector and solving the learning problem using the latent structural SVM framework [43]. To include unobserved information, Yu and Joachims [43] extended the joint feature function in structural SVM with latent variables and re-write the object function of SSVM into a difference of two convex functions. Problem of this formulation can be solved using the Concave-Convex Procedure (CCCP)[44] which is a two stages algorithm and guaranteed to converge to a local minimum.

The latent structural SVM cannot take input data directly such as structural SVM. In order to use it, inference algorithm, as well as the MRF feature function, loss function, and latent variable completion problem[43] need to be implemented first. Our implementation is described in section 4.

3 METHODS

In this section, we introduce multi-task DNNs-MRFs architecture from lower part to the top part. The whole model is constructed with two parts. The first part is a "Holistic Market Price Learner", which contains three DARNN modules. The goal of the first part is to tackle the first challenge: extracting informational representations of raw market price end-to-end without any input of hand-crafted features and technical indicators from stock market. The second part is called "Section Rotation Predictor" which is a binary Markov Random Fields model contains weighted higher order energy functions. Those higher order functions are applied to financial experts defined sector lists (used as maximum cliques). Leading stocks and lagging stocks are assigned to higher and lower weights in energy function accordingly. The goal of this part is to demonstrate second and third challenge. Unary features learned by DNNs are utilized together with higher order consistency relationships among stocks which belong to the same sector. We summarized the whole architecture in figure 1.

3.1 Holistic Market Price Learner

The "Holistic Market Price Learner(HMPL)" contains two levels, three modules of DARNNs[33]. Recall the goal of HMPL is to replace hand-crafted features and financial technical indicators with

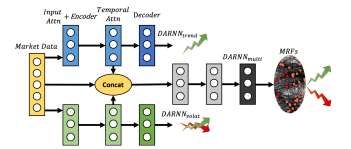


Figure 1: Multi-tasks DNN MRFs Architecture. Note that the output of DARNN $_{
m multi}$ only corresponds to one node's unary feature in MRFs.

representations learned end-to-end by neural networks. To achieve this, HMPL needs to be trained on a set of diverse and complementary tasks in order to encode holistic market price information. Bottom level contains two separate DARNN modules. They are supervised by low-level tasks such as regression to future price and volatility using raw market price data. At top level, it is supervised by high-level task that learns to use representations extracted by two bottom modules as well as raw market price data to predict ascending / descending price movement of stocks. Logits of the last layer are passed to Intra-clique Predictor described in section 3.2 as unary features.

All three DARNN modules share the same raw market price data. Here we denote the time-series dataset as X where $X = (x_1, x_2, \ldots, x_T) \in \mathbb{R}^{N \times T}$. Here $\mathbf{x}^n = (x_1^n, x_2^n, \ldots, x_T^n) \in \mathbb{R}^T$ denotes a driving series of T time-steps and $\mathbf{x}_t = (x_t^1, x_t^2)$

 $,\dots,x_t^N)\in\mathbb{R}^N$ denotes a snapshot at time-step t of all N features. For both DARNN modules at the bottom level, the input matrix is a concatenation of exogenous matrix $X\in\mathbb{R}^{5\times T}$ which contains 5 exogenous driving series: opening price, low price, high price, volume, amount and 1 target series $\boldsymbol{y}=(y_1,y_2,\dots,y_T)\in\mathbb{R}^T$. The task of those DARNNs are to predict target series y_{t+p} in the next p time steps:

$$\hat{y}_{t+p} = \text{DARNN}(y_1, \dots, y_t, x_1, \dots, x_t)$$

The target series y_{trend} of DARNN_{trend} is closing price. The target series y_{volat} of DARNN_{volat} is the standard deviation of closing price over M time-steps. We use Mean Squared Error (MSE) as loss function to train those two modules separately.

To train the top level module, which is a classification DARNN, we concatenate context vectors \boldsymbol{c}_t from each of bottom level module's second stage encoder and raw market price matrix as input matrix. The target series $\boldsymbol{y}^{\text{binary}}$ is constructed by the sign function $y_t^{\text{binary}} = sign(y_{t+p} - y_t)$ where y_t denotes closing price at time-step t. We use cross-entropy as loss function to train the final DARNN_{multi}. Logits (outputs before going through softmax) of DARNN_{multi} are then passed to "Intra-clique Predictor" as unary features.

In order to train HMPL together with MRFs in an end-to-end manner, we follow the subgradient method proposed by Witoonchart and Chongstitvatana [41]. Since our inner loop proposed in section 4.2 is actually a Structural SVM. Only gradients of parameters and feature functions need to be re-calculated. In our framework, outputs of HMPL are only used as unary features in MRFs' energy functions, our back-propagation rules can be defined by taking derivative of equation (3):

$$\frac{\partial L}{\partial \mathbf{w}^U} = \psi^U(y) - \psi^U(y^*) \tag{1}$$

where y is the ground-truth label and y^* is inferenced label. ψ^U is unary feature function described in section 3.2.1, here it contains logits calculated from DARNN_{multi}. w is parameter vector of DARNN_{multi}. And gradients of parameters can be calculated by

$$\frac{\partial L}{\partial u^U} = \mathbf{w}^U \tag{2}$$

Equations (1) and (2) can be directly plugged into sub-gradient algorithm proposed in [41]. Other configurations stay the same with their algorithm.

3.2 Intra-clique Predictor

We begin with a brief review of our choices of unary, pairwise and higher-order potential functions. We then show how to perform exact inference in models with these potentials. In section 4 we will discuss learning the parameters under the latent structural SVM framework and also how to back-prop gradients to neural networks.

3.2.1 Higer Order Energy: The Weighted Lower Linear Envelope Function. Energy functions can be decomposed over nodes N, edges \mathcal{E} and higher order cliques C [38]. Let w be vector of parameters and ψ be arbitrary feature function, then the energy can be decomposed as a set of linear combinations of weights and feature vectors:

$$E(\mathbf{y}; \mathbf{w}) = \sum_{i \in \mathcal{N}} \mathbf{w}_{i}^{U} \psi^{U}(\mathbf{y}_{i}) + \sum_{(i,j) \in \mathcal{E}} \mathbf{w}_{ij}^{P} \psi^{P}(\mathbf{y}_{i}, \mathbf{y}_{j}) + \sum_{\mathbf{y}_{C} \in \mathcal{C}} \mathbf{w}_{C}^{H} \psi^{H}(\mathbf{y}_{C})$$
(3)

where U denotes unary terms, P denotes pairwise terms and H denotes higher order terms. In this section we mainly focus on one class of higher-order potentials ψ^H defined as a concave piecewise linear function which is known as lower linear envelope potentials. This has been studied extensively in Markov Random Fields area for encouraging consistency over large cliques [12, 23, 31].

Let C denotes the set of all maximal cliques in an image and $\boldsymbol{y}_c = \{y_i | \text{ for } i \in C_j\}$ denotes set of binary random variables where $y_i \in \{0,1\}$ in clique C_j , a weighted lower linear envelope potential over \boldsymbol{y}_c is defined as the minimum over a set of K linear functions as:

$$\psi_c^H(\boldsymbol{y}_c) = \min_{k=1}^{K} \left\{ a_k W_c(\boldsymbol{y}_c) + b_k \right\}. \tag{4}$$

where $W_c(\boldsymbol{y}_c) = \sum_{i \in c} w_i y_i$ with $w_i^c \ge 0$ and $\sum_{i \in c} w_i^c = 1$ which are weights for each clique. $(a_k, b_k) \in \mathbb{R}^2$ are the linear function parameters. We illustrate an example with four linear functions in Figure 2.

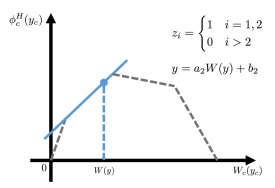


Figure 2: Example piecewise-linear concave function of $W_c(y_c) = \sum_{i \in c} w_i^c y_i$. Assume the second linear function is active namely $z^c = (1, 1, 0, 0)$ (equation 8). The result of linear combination of parameter vector and feature vector is same as quadratic psuedo-Boolean function.

Inference on energy function contains lower linear potentials is the same as the standard equation (3) and is given by:

$$\mathbf{y}^* = \operatorname{argmin} E(\mathbf{y}) \tag{5}$$

To ensure potentials do not contain redundant linear functions (functions that would never be active), Gould [13] proposed a constraint on parameters of the envelope. The *k*-th linear function is not redundant if the following condition is satisfied:

$$0 < \frac{b_k - b_{k-1}}{a_{k-1} - a_k} < \frac{b_{k+1} - b_k}{a_k - a_{k+1}} < 1.$$
 (6)

Another important property of equation (5) is shift invariant (vertically). We write $\widetilde{\psi}_c^H(\boldsymbol{y}_c)$ by shift equation (4) vertically with an abitrary amount $b^{const} \in R$

$$\widetilde{\psi}_c^H(\boldsymbol{y}_c) = \min_{k=1,\dots,K} \left\{ a_k W_c(\boldsymbol{y}_c) + b_k + b^{\text{const}} \right\}$$

Then we have

$$\underset{\boldsymbol{y}_{c}}{\operatorname{argmin}} \psi_{c}^{H}(\boldsymbol{y}_{c}) = \underset{\boldsymbol{y}_{c}}{\operatorname{argmin}} \widetilde{\psi}_{c}^{H}(\boldsymbol{y}_{c}). \tag{7}$$

Therefore, in the following discussion without loss of generality we assume $b_1 = 0$ thus $b_k \ge 0$ for k = 1, ..., n.

3.2.2 Exact Inference. Exact inference on MRFs has been extensively studied in past years. Researchers found that, energy functions which can be transformed into quadratic pseudo-Boolean functions [18, 19, 34] are able to be minimized exactly using graphcuts like algorithms [11, 15] when they satisfy submodularity condition [4]. Kohli et al. [24] and Gould [12] adapted those results to perform exact inference on lower linear envelope potentials. In this section we mainly focus on describing the st min cut graph constructed by Gould [12, 13] for exact inference (5) of energy function containing lower linear envelope potentials.

Following the approach of Kohli and Kumar [22], Gould [12, 13] transformed the weighted lower linear envelope potential (4) into a quadratic pseudo-Boolean function by introducing K-1 auxiliary variables $z=(z_1,\ldots,z_{K-1})$ with $z_k\in\{0,1\}$:

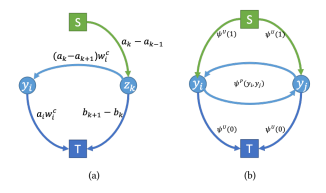


Figure 3: st-graph construction for equation (9), unary and pairwise terms. Every cut corresponds to an assignment to the random variables, where variables associated with nodes in the S set take the value one, and those associated with nodes in the T set take the value zero. With slight abuse of notation, we use the variables to denote nodes in our graph.

$$E^{c}(\boldsymbol{y}_{c}, z) = a_{1}W_{c}(\boldsymbol{y}_{c}) + b_{1} + \sum_{k=1}^{K-1} z_{k} \left((a_{k+1} - a_{k}) W_{c}(\boldsymbol{y}_{c}) + b_{k+1} - b_{k} \right)$$
(8)

for a single clique $c \in C$. Under this formulation, minimizing the pseudo-Boolean function over z is equivalent to selecting (one of) the active functions(s) from equation (4). Another important property of optimized z under this formulation is that it automatically satisfies the constraint: $z_{k+1} \leq z_k$. This property give rise to further development of parameter vector and feature vector (equation (12) and (13)) which are used in latent structural SVM.

In order to construct the st-min-cut graph, we rewrote equation (8) into posiform [4]:

$$E^{c}(\boldsymbol{y}_{c}, \boldsymbol{z}) = b_{1} - (a_{1} - a_{K}) + \sum_{i \in c} a_{1} w_{i}^{c} y_{i}$$

$$+ \sum_{k=1}^{K-1} (b_{k+1} - b_{k}) z_{k} + \sum_{k=1}^{K-1} (a_{k} - a_{k+1}) \bar{z}_{k}$$

$$+ \sum_{k=1}^{K-1} \sum_{i \in c} (a_{k} - a_{k+1}) w_{i}^{c} \bar{y}_{i} z_{k}$$

$$(9)$$

where $\bar{z}_k = 1 - z_k$ and $\bar{y}_i = 1 - y_i$. a_1 is assumed to be greater than 0 so that all coefficients are positive (recall we assume $b_1 = 0$ in section 3.2.1 and we have $a_k > a_{k+1}$ and $b_k < b_{k+1}$). Since the energy function (9) is submodular, the *st-min-cut* graph can be constructed based on equation (9). The construction (including unary and pairwise) is explained in Figure 3.

4 OPTIMIZATION

4.1 Transforming Between Representations

With the inference algorithm in hand, we now can develop the learning algorithm for weighted lower linear envelope potentials using the latent structural SVM framework. We begin by transforming the equation (8) into a linear combination of parameter vector and feature vector. Then a two-step algorithm was developed to solve the latent structural SVM.

The latent structural SVM formulation requires that the energy function be formulated into a linear combination of features and weights while our higher-order potential is represented as the minimum over a set of linear functions. However, in 3.2.2 we reformulated the piesewise linear functions into a quadratic pseudo-Boolean function (8) by introducing auxiliary variables. Now we show function (8) itself is an inner product of parameter vector and feature vector with latent information. Note that the function can be expanded as a summation of 2K-1 terms:

$$E^{c}(y_{c}, z) = a_{1}W_{c}(y_{c}) + b_{1}$$

$$+ \sum_{k=1}^{K-1} z_{k}((a_{k+1} - a_{k})W_{c}(y_{c}) + b_{k+1} - b_{k})$$

$$= a_{1}W_{c}(y_{c}) + \sum_{k=1}^{K-1} (a_{k+1} - a_{k})z_{k}W_{c}(y_{c})$$

$$+ \sum_{k=1}^{K-1} (b_{k+1} - b_{k})z_{k}$$
(10)

Here we use the fact of equation (7) and let $b_1 = 0$. Now we can reparameterize the energy function as

$$E^{c}(\boldsymbol{y}_{c}, z; \boldsymbol{\theta}) = \boldsymbol{\theta}^{T} \psi(\boldsymbol{y}_{c}, z)$$
(11)

where:

$$\theta_k = \begin{cases} a_1 & \text{for } k = 1\\ a_k - a_{k-1} & \text{for } 1 < k \le K\\ b_{k+1-K} - b_{k-K} & \text{for } K < k \le 2K - 1 \end{cases}$$
 (12)

$$\psi_{k} = \begin{cases} W_{c}(\boldsymbol{y}_{c}) & \text{for } k = 1\\ W_{c}(\boldsymbol{y}_{c}) z_{k} & \text{for } 1 < k \le K \\ z_{k} & \text{for } K < k \le 2K - 1 \end{cases}$$
(13)

Under this formulation, inference problems in [43] can be written as:

$$(\hat{\mathbf{y}}_{k}(\boldsymbol{\theta}), \hat{\mathbf{z}}_{k}(\boldsymbol{\theta})) = \underset{(\mathbf{y} \times \mathbf{z}) \in \mathcal{Y} \times \mathcal{Z}}{\operatorname{argmin}} \boldsymbol{\theta}^{T} \cdot \psi(\mathbf{y}_{k}, \mathbf{z}_{k})$$
(14)

and

$$\mathbf{z}_{k}^{*}(\boldsymbol{\theta}) = \operatorname*{argmin}_{\mathbf{z} \in \mathcal{T}} \boldsymbol{\theta}^{T} \cdot \psi(\mathbf{y}_{k}, \mathbf{z}_{k}) \tag{15}$$

There are 2 facts worth to mention. The first fact is that in our previous construction of minimum-st-cut graph the latent variable z is already included. Therefore, we can apply our inference algorithm directly on our 2 new formulations.

More interesting for equation (15) there exists more efficient algorithm. At training stage the ground-truth labels y_i is a function input thus completely observed. Therefore, the term ($(a_{k+1} - a_{k+1})$)

 $a_k)W_c(\boldsymbol{y}_c)+b_{k+1}-b_k)$ in equation (10) becomes constant. So we can infer latent variable z explicitly by:

$$z_k^c = \begin{cases} 0 & \text{if } ((a_{k+1} - a_k)W_c(y_c) + b_{k+1} - b_k) \ge 0\\ 1 & \text{otherwise.} \end{cases}$$
 (16)

Therefore, assignments inferred by graph-cut algorithm can be directly encoded into a linear combination by using our latent structural SVM formulation for learning purpose. The remaining task is to ensure the concavity of θ . We do this by adding following constraint:

$$A\theta \ge \epsilon, \ A = \begin{bmatrix} 1 & 0 & 0 \\ 0 & -1 & 0 \\ 0 & 0 & P \end{bmatrix} \in \mathbb{R}^{(2K-1)\times(2K-1)}$$
 (17)

where -1 is a matrix of size $(K-1)\times (K-1)$ and **P** is an identity matrix of size $(K-1)\times (K-1)$. One subtle problem we found during experiments is that the algorithm can be stuck with small numerical value. To avoid this we add small slack variables $\epsilon = \mathbf{1}^{-15}$ on those constraints.

4.2 Latent Structural SVM Learning

Algorithm 1 Learning lower linear envelope MRFs with latent variables.

```
1: Set MaxIter = 100
 2: input training set \{y_i\}_{i=1}^n, regularization constant C > 0, and
      tolerance \epsilon \geq 0
 3: Initialize \theta using Algorithm 2
 4: repeat
         CCCP Outer Loop
 5:
         Set iter = 0
 6:
         for each training example, i = 1, ..., n do
 7:
             compute z_i^* = \operatorname{argmax}_{\mathbf{z} \in \mathcal{Z}} \theta \cdot \psi(\mathbf{y}_i, \mathbf{z})
 8:
 9:
         initialize active constraints set C_i = \{\} for all i
10:
         repeat
11:
             CCCP Inner Loop
12:
13:
             solve the quadratic programming problem in equation 20
             with respect to active constraints set C_i for all i and con-
             cavity constraints A\theta \geq \epsilon to get \hat{\theta} and \hat{\xi}
             for each training example, i = 1, ..., n do
14:
                 compute \hat{\boldsymbol{y}}_i, \hat{\boldsymbol{z}}_i = \operatorname{argmin}_{\boldsymbol{y}} E(\boldsymbol{y}, \boldsymbol{z}; \hat{\boldsymbol{\theta}}) - \Delta(\boldsymbol{y}, \boldsymbol{z}, \boldsymbol{y}_i)
15:
                 if \hat{\xi}_i + \epsilon < \Delta(\hat{y}_i, \hat{z}_i, y_i) - E(\hat{y}_i, \hat{z}_i; \hat{\theta}) + E(y_i, z_i^*; \hat{\theta}) then
16:
                     C_i \leftarrow C_i \cup \{\boldsymbol{y}_i^{\star}\}
17:
                 end if
18:
             end for
19:
         until no more violated constraints
20:
         return parameters \hat{\theta}
21:
         Set iter = iter + 1
22:
23: until iter \ge MaxIter
24: return parameters \hat{\boldsymbol{\theta}}
```

With the inner product formulation (equation (11)) of higher order energy function in hand, we now able to develop our latent structural SVM learning algorithm. The energy function (higher

order function together with unary and pairwise functions) can be written as:

$$E_{all}(y,z) = \begin{bmatrix} \theta^{H} \\ \theta^{unary} \\ \theta^{pairwise} \end{bmatrix}^{T} \cdot \begin{bmatrix} \psi^{H} \\ \psi^{unary} \\ \psi^{pairwise} \end{bmatrix} = \theta^{T}_{all} \cdot \psi_{all}$$
 (18)

where $\theta^H \in \mathbb{R}$ is the parameter vector in higher order equation (11) of size 2K-1. θ^{unary} and $\theta^{pairwise}$ are both scalars. $\psi^{unary} = \sum_i \psi_i^U(y_i)$ and $\psi^{pairwise} = \sum_{ij} \psi_{ij}^P(y_i, y_j)$. Therefore, the size of θ_{all} is 2K+1.

Following Yu and Joachims [43], we use the two stages Concave-Convex Procedure (CCCP) [44] to solve the optimization problem. We first imputes the latent variables *z* explicitly by equation (15). Namely solving the "latent variable completion" problem [43]:

$$\mathbf{z}_{i}^{*} = \underset{\mathbf{z} \in \mathcal{Z}}{\operatorname{argmax}} \, \theta \cdot \psi(\mathbf{y}_{i}, \mathbf{z}) \tag{19}$$

The inference result z_i^* for $i=1,\ldots,n$ is used as completely observed for later stage. With the latent variable z_i^* which best explains the ground-truth data y_i in hand, updating the parameter vector $\boldsymbol{\theta}$ reduces to solve the standard structural SVM problem:

$$\min_{\theta} \left(\frac{1}{2} \|\theta\|^2 + C \sum_{i=1}^{n} \left(\max_{(\hat{\mathbf{y}} \times \hat{\mathbf{z}}) \in \mathcal{Y} \times \mathcal{Z}} [\theta \cdot \psi(\hat{\mathbf{y}}, \hat{\mathbf{z}}) + \Delta(\mathbf{y}_i, \hat{\mathbf{y}}, \hat{\mathbf{z}})] \right) \right) (20)$$

$$-C \sum_{i=1}^{n} \left(\theta \cdot \psi(\mathbf{y}_i, \mathbf{z}_i^*) \right)$$

Our optimization algorithm is summarized in Algorithm 1. As we mentioned in Appendix A, although we proposed an end-to-end subgradient algorithm is section 3.1, MRFs updated by such algorithm take too many iterations to converge. Therefore, we propose a two-stage training procedure. At first stage, HMPL and MRFs are trained separately. Therefore, MRFs can take advantage of the efficient latent structural SVM and converge in a polynomial number of iterations. After all those models are converged, we then combine them together to conduct end-to-end training. Note that the CCCP Inner Loop in Algorithm 1 is actually solving standard structural SVM problem. Therefore, at the second stage, we use subgradient algorithm proposed in section 3.1 to replace the CCCP Inner Loop. Other settings remain the same.

The last problem remaining is the initialization method. Because our objective function (20) is not convex and the CCCP algorithm is only guaranteed to converge to a local minimum or saddle point [44], initialization of θ might affect the performance of our algorithm. Since there are no theoretical solution for this problem, we propose an empirical initialization algorithm in Appendix A.1.

5 EXPERIMENT

In this section, we first introduce 3 stock datasets (index) and how we use them to construct our final input datasets. Then, we introduce the parameter settings of our model and other training details. Finally, we select four evaluation metrics and use them to demonstrate our framework's effectiveness by comparing with several baseline methods.

5.1 Dataset and Model Settings

To demonstrate the effectiveness of higher order consistency, we choose three exclusive and the most famous stock indexes on Chinese stock market to build our input datasets. Their index codes are: CSI (China Securities Index) 200, CSI 300 and CSI 500 which contain 200, 500 and 300 constituent stocks respectively. The CSI 300 index selects most liquid A-share stocks. It aims to reflect the overall performance of China A-share market. The CSI 200 and 500 index aims to reflect the overall performance of mid-to-large and small-to-mid capital A-shares respectively.

All these indexes are exclusive and are refined on a yearly basis. In this paper, we use fixed versions on 30-JAN-2015. We then collect their constituent stocks' minute-level data from 05-JAN-2015 to 29-DEC-2017. On Chinese stock market each trading day has 3 trading hours. So there are 240 samples for each normally traded stock each day. Each sample contains 6 features: opening price, high price, low price, closing price, volume, amount. ¹ For each stock, the first 80% days are used to construct the training set and the last 20% days are used as testing set. Approximately training set and testing set contain 13 million and 3 million samples, respectively.

Table 1: Technical Indicators Selection

Category	Indicator Name						
Momentum	Awesome Oscillator, Money Flow Index						
Volume	Chaikin Money Flow						
	On-balance volume mean						
Volatility	Bollinger Bands (Upper and Lower Bands)						
Trend	Average Directional Movement Index						
	Moving Average Convergence Divergence						

To demonstrate benefits of multi-task RNN over manually designed technical indicators, we also need to construct technical indicators datasets for each of those market price dataset collected above. We select 8 most popular indicators, 2 from each category [21] shown in Table 1. In implementation, we use open source package *Technical Analysis Library in Python*² to calculate those indicators and all hyper parameters are using package's default settings. After technical indicators calculation, these 8 new features are concatenated to above market price dataset (5 features at each minute). So the final input dataset for each single task model contains 13 features in total. Before feeding into models, we normalize each stock with *z-score* function using standard deviation and mean calculated in the training set.

For brevity, we denote market price dataset which only contains 5 features as **Market** and the concatenated 13 features dataset as **Indicator**. As discussed in section 3.1, closing price at time t can be directly used as regression target for DARNN $_{\rm trend}$. Standard deviation of closing price with a window size of 10 is used as regression target for DARNN $_{\rm volat}$. The dimensions of hidden state and cell state are 32 for DARNN $_{\rm trend}$ as well as DARNN $_{\rm volat}$ and 128

for $\mathsf{DARNN}_{\mathsf{multi}}.$ More training details are described in appendix A.

5.2 Results

In order to demonstrate the effectiveness of our framework, we run 3 baselines on 3 different Chinese Securities Indexes with and without technical analysis indicators as inputs. Results are summarized in Table 2. All results are reported over the test sets. We select four metrics as evaluation metrics to justify the effectiveness of the proposed approach. They are calculated by collecting all predicted labels of constituent stocks in each CSI index.

5.2.1 Effectiveness of multi-task framework. As mentioned earlier, to demonstrate effectiveness of multi-task framework, we use Indicator dataset, which contains both market price data and technical analysis indicators as inputs for DARNN and Market dataset which only contains market price data as inputs for HMPL and other single task models. For vanilla DARNN, we use a hidden size of 128. HMPL's configuration is described in section A.2. As we can see in Table 2, single task models (LSTM, LSTM_ATTN, DARNN) tested on Market dataset (without technical analysis indicators as inputs) generally have worse performance on all 4 metrics. In particular, performance of DARNN models tested on Indicator dataset is consistently better than the ones on Market dataset. This proves that even with hand-crafted features, deep learning models can still benefit from diversified and complementary features. By comparing DARNN tested on Indicator and HMPL on Market, we can see that HMPL marginally outperforms DARNN on CSI200 and CSI300 index constituent stocks and is slightly worse on CSI500 constituent stocks. We can conclude that by using multi-task RNN, we can extract equally good features compared with hand-crafted features.

5.2.2 Effectiveness of higher-order MRFs. In Table 2, we can observe that HMPL-MRFs framework consistently outperforms other baselines on all 3 CSI index constituent stocks. It shows evidence that higher-order energy function can help with encoding clique level consistency thus improving overall prediction performance. One interesting point to note is that the recall rate of HMPL-MRFs is constantly lower than other baselines. This can be seen as a trade off between accuracy and recall rate. However, it is worth to mention that for stock price movement prediction, high accuracy and precision are much preferred than recall rate. Another interesting phenomenon is that HMPL-MRFs gives higher improvements on CSI200 and CSI300 while little improvements over DARNN trained with technical analysis indicators on CSI500. One possible reason is that CSI200 and CSI300 select most liquid and representative stocks in Chinese stock market. Those stocks exhibit much stronger and higher order consistency than illiquid stocks. CSI500 selects smallmid capital stocks which are less liquid and contains much more noisy movements.

5.2.3 Visualization of higher-order consistency. In order to further investigate higher-order MRFs' effectiveness, we design a heat-map to visualize CSI300 index intra-clique higher-order relationship in figure 4.

 $^{^1\}mathrm{During}$ this period, there are some stocks de-listed (SZ000024, SH600485, SH600832 in CSI 200; SZ000693, SZ000748, SZ000982 in CSI 500; SH600485, SH600832, SZ000024, SH601299 in CSI 300). Therefore, in total we collect 197, 497 and 296 stocks during this period respectively.

²https://github.com/bukosabino/ta

Table 2: Results: Baselines and ablation study. All models have a window size (lag steps) of 20 and predict price movement label at the next time step.

Data Set	Models		Chinese Securities Index (CSI)											
			CSI200				CSI500				CSI300			
		Accuracy	Precision	Recall	F1 Score	Accuracy	Precision	Recall	F1 Score	Accuracy	Precision	Recall	F1 Score	
	LSTM	62.30	73.82	70.70	72.23	60.35	68.56	70.03	69.29	60.16	71.39	68.50	69.91	
Indicator	LSTM_ATTN	64.26	72.41	74.34	73.36	61.13	75.63	66.67	70.86	64.26	75.54	70.57	72.97	
	DARNN	63.09	72.08	73.55	72.81	66.60	78.98	74.13	76.48	65.82	76.68	73.46	75.04	
Market	LSTM	57.62	67.57	67.37	67.47	55.86	68.10	64.53	66.27	56.25	67.17	65.98	66.57	
	LSTM_ATTN	59.57	71.60	66.86	69.15	58.40	69.53	68.12	68.81	61.33	71.87	68.91	70.36	
	DARNN	61.13	71.26	71.47	71.37	63.09	77.04	67.87	72.16	63.87	72.09	73.59	72.83	
	HMPL	65.04	74.04	73.39	73.72	65.43	76.04	72.80	74.38	66.60	71.67	78.90	75.11	
	HMPL+MRFs	67.97	77.51	73.91	75.67	66.80	79.65	72.78	76.06	68.95	78.55	74.71	76.58	
				A Company of the Control of the Cont				08 04 00 -04			ì			
(a) Ground-truth					(b) HMPL						(c) HMPL-MRFs			

Figure 4: Higher order consistency visualization. Figure (a) is calculated directly from ground truth labels on test set. Figure (b) is calculated using predicted labels of HMPL without MRFs on the test set. In figure (c), we use predicted labels of HMPL-MRFs on test set as inputs.

We first select two sectors: nonferrous metal sector, which contains 10 constituent stocks, and infrastructure sector, which contains 35 constituent stocks from CSI300 index 3 . We then measure consistency level between each two of these constituent stocks. In order to capture their temporal relationship, we propose a novel consistency measure which is calculated on temporal intervals.

Let $\boldsymbol{y}_i^T = \{y_i^1, y_i^2, ..., y_i^T\}$ denotes time-series for stock i. $y_i^t \in \{0, 1\}$ is the binary price movement label at time t. We segment time-series \boldsymbol{y}_i^T into $N = \lceil \frac{T}{P} \rceil$ non-overlapping intervals $\{y_i^n, y_i^{n+1}, ..., y_i^{n+P}\}$ with fixed length P. For any two stocks i and j, we calculate the difference $d_{ij}^n = \sum_n^{n+P} y_i^n - \sum_n^{n+P} y_j^n$ of how many times positive price movement happen in the nth time interval in each stock. Then the consistency level c_{ij} between stocks i and j can be calculated as a ℓ_1 norm:

$$c_{ij} = -\|\mathbf{d}_{ij}\|_1$$

where $\mathbf{d}_{ij} = \{d_{ij}^1, d_{ij}^2, ..., d_{ij}^N\}$. We normalize c_{ij} into interval [-1, 1]. Each entry in figure 4 denotes a consistency level measure c_{ij} . The larger the c_{ij} is, the higher of consistency level between stock i and stock j, the color of corresponding entry is closer to red, and vice

versa. As we mentioned, the average duration of information arrival-conduction-integration-release process is 4.04 minutes. Since which stock is leading at each time interval is elusive, we set P=9 when calculating consistency measures.

As we can see in figure (a), there is a significant red square area, which means ground-truth heat-map shows strong intra-clique consistency. This is an evidence that higher-order relationships do exist within clique of stocks. However, in figure (b), the red square area is fragmented into many little pieces. The whole area's color is closer to blue when compared to ground-truth heat-map, which means that HMPL captures little higher-order consistency. The reason we still can observe a shape of red square is that the accuracy of HMPL model on CSI300 is 66.6%. However, we can still conclude that the accuracy of single HMPL model mainly comes from unary features and it fails to capture higher order consistency relationships lies in clique of stocks. On the contrary, even though HMPL-MRFs model's accuracy on CSI300 index is only 2.35% better than HMPL model, we can observe that heat-map (c) is more close to ground-truth heat-map than heat-map (b). There is a much clear red square and the number of small fragments in red area is also less than figure (b). We can conclude that HMPL-MRFs models learn to utilize both unary features from HMPL as well as higher-order relationships encoded in MRFs.

³These two sectors are selected only because painting many sectors in one figure would be too messy to interpret and those two sectors have appropriate clique size (number of stocks) for visualization. Conclusions from these two sectors also apply to other sectors

6 CONCLUSION

This paper has shown how to model individual stock price prediction problem without hand-crafted features and encode lead-lag relationships among stocks using weighted higher-order MRFs. A multi-task neural networks framework, Holistic Market Price Learner (HMPL) is proposed to automatically extract diversified and complementary features from individual stock price sequence. Features learned by HMPL are passed to a binary MRFs with weighted lower linear envelope energy function to utilizing intra-clique higher order consistency among stocks. An efficient latent structural SVM algorithm is designed for learning MRFs in polynomial time. Finally the MRFs and HMPL are trained end-to-end using subgradient algorithm. Extensive experiments are conducted on three major index on Chinese stock market. The proposed HMPL-MRFs achieves best accuracy on all three indexes.

Our work gives insights to a number of directions for future research. One obvious extension is to apply our method to a multilabel MRFs to help with 3-classes price movement prediction. A More interesting direction is to investigate the implicit relationship between the expert defined index list and Graph RNN [42]. Investigating under this direction could help further diminishing domain knowledge required by our framework.

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A TRAINING DETAILS

A.1 Initialization of lower linear envelope

We assume that the more evenly distributed of $W_c(Y_c)$ where $c \in C$ on x axis, the more rich representation (number of linear functions) the energy function should have. In order to initialize θ , we first determine the x-coordinate of sampled points sp. Then we sample its y-coordinate from a uniform distribution $\mathcal{U}(\text{upbound}, \text{upbound}-0.5)$ to add some randomness in our initialization as well as maintain concavity. Linear parameters a_k and b_k are later calculated using those sampled points sp_k and sp_{k-1} . At last we encode $\{a_k, b_k\}_{k=1}^K$ into θ using equation (12). This algorithm is summarized in Algorithm 2.

Algorithm 2 Empirical initialization algorithm for θ

```
1: gap = \frac{1}{K}, a_1 = \mathcal{U}(0, 1e6), b_1 = 0, sp_1 = (0, 0), w_0 = 0, counter = 2
2: for each clique c \in C do
3: Compute weighted clique value w_c = W_c(y_C)
4: if w_c - w_{c-1} > gap then
5: upbound = a_{counter}w_c + b_{counter} sp_{counter} = (w_c, \mathcal{U}(upbound - 0.5, upbound)) Calculate a_{counter} and b_{counter} using sp_{counter-1} and sp_{counter} counter = counter + 1
6: end if
7: end for
8: If counter < K, remaining a and b are all set to be a_{counter} and a and a and a are all set to be a and a and a are all set to be a and a are all set to be a and a and a are all set to be a and a and a are all set to be a are all set to be a and a are all set to be a are all set to be a and a are all set to be a are all set to be a and a are all set to be a and a are all set a and a are all set a are all set a and a are all set a are all set a and a are all set a and a are all set a and a are
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A.2 Multi-task training

To improve accuracy and reduce over-fitting, we add a drop out layer between input layer and LSTM layer with a ratio of 0.2. We also clip and normalize gradients during back-propagation stage with a maximum norm of 5.0 to prevent gradient exploding issue. As pointed out by Lample et al. [26], the question of "when should the training schedule switch from one task to another task?" or "should each task be weighted equally?" remains open. In our implementation, we follow the proportional sampling approach described by Søgaard and Goldberg [36]. After a backward pass completed, we randomly sample a new task as well as its batch data as the next task to be trained. In practice, we use a proportion of [0.25, 0.25, 0.5] for three tasks respectively. This mechanism helps multi-task model to avoid *Catastrophic Forgetting* phenomenon which means lower level model forgets learned knowledge during higher level model back-propagation pass.

Even though we propose an end-to-end training algorithm for HMPL and MRFs in section 3.1, MRFs inference stage is still too slow to be trained jointly with HMPL. To overcome this difficulty, we implement a two stages training procedure. We first add a *softmax* layer on top of DARNN_{class} and train HMPL separately from MRFs. We use *Negative Log-likelihood* as the loss function. At the second stage, after HMPL converge, we remove the *softmax* layer and retrain it together with MRFs. One issue we must mention is that,

even though we use binary MRFs which can only predict positive / negative price movement, we find there is a significant amount of time when stock price remains no change. We find it benefits the performance a lot if we treat the classification as a three classes problem rather than a binary classification problem during the first stage. Therefore, at the first stage, the *softmax* layer will output probability for three labels: *negative movement*, *no changes* and *positive movement*. Since binary MRFs still needs a two dimension input as part of unary energy function, after the *softmax* layer is removed, we add an additional linear mapping layer between logits of HMPL and MRFs at the second stage.

A.3 End-to-end NN-MRFs training

With converged HMPL and MRFs at hand, now we can go forward to train them in an end-to-end manner. We only include pairwise energy function through section 3.2 and section 4 to show a general application of our proposed algorithm. In the case of Chinese stock market, to our best knowledge there is no public available definition of pairwise relationship between stocks. Therefore, in our implementation we only use unary and higher order energy function. Each stock is then treated as a node in MRFs and each stocks group which has lead-lag relationships is treated as a maximum clique in MRFs. One benefit of MRFs clique is that we can embed domain expert knowledge about industry classification as maximum cliques into our model. We choose to use Tonghuashun industry classification [1] in our model. One subtle but crucial detail about modeling lead-lag effect lies in equation (4). Recall that $W_c(\boldsymbol{y}_c) = \sum_{i \in c} w_i y_i$ with $w_i^c \ge 0$ and $\sum_{i \in c} w_i^c = 1$ which are weights for stocks in each clique. Therefore, leading stocks should have a higher weights while lagging stocks should have lower weights. In our implementation, we use constituents' weight defined in CSI200, CSI500 and CSI300 as their weights in equation (4) and normalize them to ensure the summation equals 1.