

# Automated band annotation for capillary electrophoresis based high-throughput RNA structure probing

## ABSTRACT

### Motivation:

### Results:

**Availability:** The implementation of the proposed algorithm is included in the HiTRACE software freely available for download at <http://hitrace.stanford.edu>.

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## 1 INTRODUCTION

RNA plays diverse roles in encoding and regulating genetic information. We can better understand this versatility of RNA by knowing its higher order structure. To this end, chemical probing methodologies provide a powerful means to mapping RNA secondary and tertiary structure at single-nucleotide resolution (Weeks, 2010).

There exist many chemical probing techniques, most of which have common experimental procedures as follows: Treated with an RNA of interest, a chemical reagent modifies the RNA, either cleaving it or forming a covalent adduct with it. Examples of the reagent include hydroxyl radicals, dimethyl sulfate (DMS), CMCT, kethoxal and bisulfite. Subsequent reverse transcription detects the modification sites as stops to primer extension at nucleotide resolution. Resulting cDNA fragments are resolved in sequencing gels followed by individually quantifying band intensities. Typically, the bottleneck is the final steps (gel running and band quantification).

To resolve fragments in high-throughput fashion, capillary electrophoresis (CE) can be used. CE-based chemical probing can easily produce tens of thousands of individual electrophoretic bands from a single experiment, leading to recent breakthroughs in high-throughput mapping of complex RNA structures (Mitra *et al.*, 2008; Vasa *et al.*, 2008; Weeks, 2010; Das *et al.*, 2010; Kladwang and Das, 2010) such as ribosomes (Deigan *et al.*, 2009), and viruses (Wilkinson *et al.*, 2008; Watts *et al.*, 2009).

Analyzing a large number of electrophoretic traces from a high-throughput structure-mapping experiment is time-consuming and poses a significant informatic challenge, requiring a set of robust signal-processing algorithms for accurate quantification of the structural information embedded in the noisy traces. Current software methods for CE analysis include capillary automated footprinting analysis (CAFA; Mitra *et al.*, 2008), ShapeFinder (Vasa *et al.*, 2008), high-throughput robust analysis for capillary electrophoresis (HiTRACE; Yoon *et al.*, 2011), and fast analysis of SHAPE traces (FAST; Pang *et al.*, 2011).

A typical high-throughput CE analysis pipeline consists of the following steps (Yoon *et al.*, 2011): preprocessing such as normalization and baseline adjustment, alignment, peak detection,

band annotation, and peak fitting. Among these, band annotation refers to the process of mapping each band in a electrophoretic trace to a position in the nucleic acid sequence. For verification, visual inspection in this phase is normally inevitable to certain extent. However, in practice, this band annotation step often takes significant human efforts in CAFA and ShapeFinder, for they were designed to focus more on alignment and peak fitting. HiTRACE and FAST provides an improved level of band annotation support, but band annotation remains still the most time-consuming step of a HiTRACE- or FAST-based analysis pipeline.

This paper describes a dynamic-programming based approach to automated band annotation for high-throughput capillary electrophoresis. Figure 1 shows the overview of the proposed methodology for automated band annotation.

## 2 METHODS

### 2.1 Experimental setup

### 2.2 Problem definition

Given an RNA sequence  $s$  of length  $N$ , assume that we carry out the chemical structure probing of this sequence using  $M$  different treatments, each of which is run by a separate capillary lane. Assume that the fluorescence intensity of each capillary is measured over  $K$  time points. We define a *profile* as the sequence of intensity values from a capillary. The entire CE measurement can then be arranged in a  $K \times M$  matrix  $\mathbf{D}$ . Normally,  $N \ll K$ . Based on the characteristic of the chemical agent used in each treatment and the secondary structure computationally inferred from the input sequence, we can predict the fluorescence intensity at each position of  $s$  for each of  $M$  treatments. This prediction can be arranged in a  $N \times M$  matrix  $\mathbf{P}$  called the *prediction matrix*.

The problem of band annotation is formulated as selecting  $N$  out of the  $K$  rows of  $\mathbf{D}$  using the information in  $\mathbf{P}$  in such a way that a certain objective is optimized over all possible  $\binom{K}{N}$  possibilities. The selected  $N$  points map to the locations of the nucleotides of the sequence  $s$  in the CE measurement.

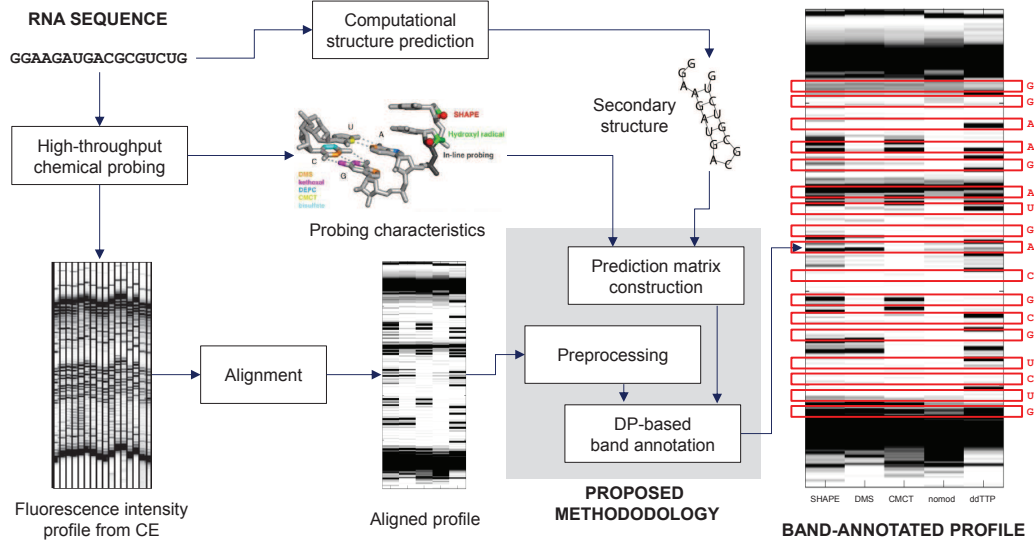
The input of the proposed method consists of the following:

- $\mathbf{D} \in \mathbb{R}^{K \times M}$ : the fluorescence intensity matrix
- $\mathbf{P} \in \{0, 1\}^{N \times M}$ : the prediction matrix
- $s \in \{A, C, G, U\}^N$ : the nucleotide sequence

and the output is an array  $\mathbf{y} \in \mathbb{Z}_+^N$  representing  $N$  points selected out of  $K$ .

### 2.3 Prediction matrix construction

Given an RNA sequence, we computationally predict its secondary structure using the Vienna RNA package (Hofacker, 2003). Based on the predicted structure and the properties of the chemical probing used, we construct the prediction matrix  $\mathbf{P}$  that stores the expected chemical reactivity for individual residues. The element  $p_{ij} \in \mathbf{P}$  indicates such reactivity information of residue  $i$  to reagent  $j$ .



**Fig. 1.** Overview of the proposed dynamic-programming-based band annotation methodology. Given an RNA sequence, we carry out high-throughput structure-mapping experiments, producing a number of capillary electrophoresis (CE) profiles. Additionally, we computationally predict the secondary structure of the input sequence. From the predicted structure and the characteristics of the chemical probing method used, we drive a prediction matrix that stores expected interaction patterns between the residues. Based on the aligned CE profiles and prediction matrix, we apply a dynamic-programming approach that finds the optimal selection of the band locations under the predetermined scoring scheme.

We assume the use of three chemical probing strategies in this paper: dimethyl sulfate alkylation (Tijerina *et al.*, 2007) [DMS], carbodiimide modification (Walczak *et al.*, 1996) [CMCT], and 2'-OH acylation [the SHAPE strategy (Merino *et al.*, 2005)]. Figure 2(a) defines the expected reactivity of each type of nucleotide to chemical reagents used for chemical probing under the (un)paired condition. The value of one means the reactivity to a reagent (*i.e.*, the existence of a band in the fluorescence profile), whereas zero indicates no reactivity (*i.e.*, no band). For instance, in the unpaired condition, the DMS chemical modifies A and C but not U and G, and the entries for A and C are one, while those for U and G are zero.

Figure 2(b) shows an example RNA sequence with its secondary structure. Figure 2(c) shows the corresponding prediction matrix  $\mathbf{P}$ .

## 2.4 Preprocessing intensity data

Let  $\mathbf{d}_j$  be the  $j$ -th column vector of  $\mathbf{D}$ ,  $1 \leq j \leq M$ . The following procedure is executed.

1. Select candidates for the peaks in  $\mathbf{d}_j$  that can be mapped into elements of the sequence  $\mathbf{s}$ . These peaks are selected to satisfy the following conditions. First, a peak  $\mathbf{d}_j(k)$  must have a higher intensity (a fundamental property of a peak) than those of its neighbors,  $\mathbf{d}_j(k-1)$  and  $\mathbf{d}_j(k+1)$ . Second, a peak must have a high negative gamma which is roughly estimated as  $\delta^+ - \delta^-$ , where  $\delta^- = \max(\mathbf{d}_j(k) - \mathbf{d}_j(k-1), (\mathbf{d}_j(k) - \mathbf{d}_j(k-2))/2)$  and  $\delta^+$  is defined similarly. We choose  $n$  peaks with highest negative gammas from the points satisfying the first condition, while  $n$  may vary according to the data set type.
2. In order to remove the influence of noise that every  $\mathbf{d}_j$  has in common near the end of the time series, eliminate a portion of tailing peaks as follows: Identify 5 tailing peaks and select one with the highest intensity among them. Denote  $i_j$  be the time index of this peak ( $1 \leq i_j \leq K$ ). Let  $i^* = \max_{1 \leq j \leq M} i_j$ . For each  $\mathbf{d}_j$ , remove all peaks appearing after  $i^*$ .
3. Based on the remaining peak locations, construct a matrix called the *bonus matrix*  $\mathbf{B} \in \mathbb{Z}^{K \times M}$ . Let  $-\bar{\gamma}$  be the mean value of negative

gamma of the remaining peaks. Initialize  $\mathbf{B}$  to all zero. If  $\mathbf{D}(i, j)$  represents a peak, then we set  $\mathbf{B}$  to be the negative gamma of the corresponding peak added by  $-\bar{\gamma}/2$ .

4. Determine the ideal separation between bands based on the remaining peak locations:  $\rho \triangleq (k_r - k_f)/(N - 1)$ , where  $k_f$  and  $k_r$  are the locations of the foremost peak and the rearmost peak respectively. The ideal separation calculated here is used when shifting the window in dynamic programming.

## 2.5 Formulation as dynamic programming

In essence, the band annotation problem is to select  $N$  out of  $K$  points and match them to peak locations - if at all possible - in an optimal way. This is similar to the problem of aligning two sequences  $(1, 2, \dots, N)$  and  $(1, 2, \dots, K)$  without allowing gaps for the latter. With regard to the problem of mapping  $K$  points into peak locations, the relative location of the corresponding peak is assigned to each element of the former sequence as shown in the following example:

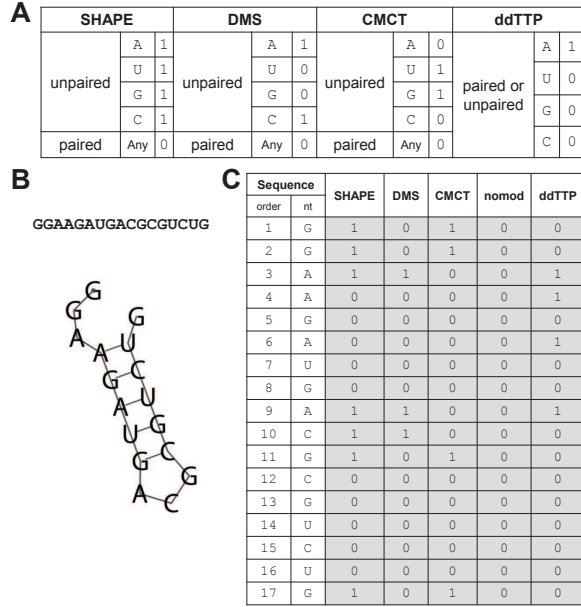
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RNA sequence index : -1--2---3...N...-
Measurement index : 123456789.....K
Relative peak location: -2--0---1...0...-

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The first three bands are located at 2, 5, and 9 time units. The relative peak location assigned to the third band is 1 implying that this band corresponds to the peak located 1 unit behind it, namely at 10 time unit. Each set of alignment and peak location correspondence is given a score that represents its probabilistic likelihood. Dynamic programming can be utilized to find the solution set with the highest score, which in turn leads to the most likely locations of bands.

One difficulty that arises in the dynamic programming approach is that the number of possibilities to be considered grows large as the number of lanes increases; if there are  $M$  lanes to consider,  $M + 2$  sequences must be taken into account for alignment in the dynamic programming, resulting in



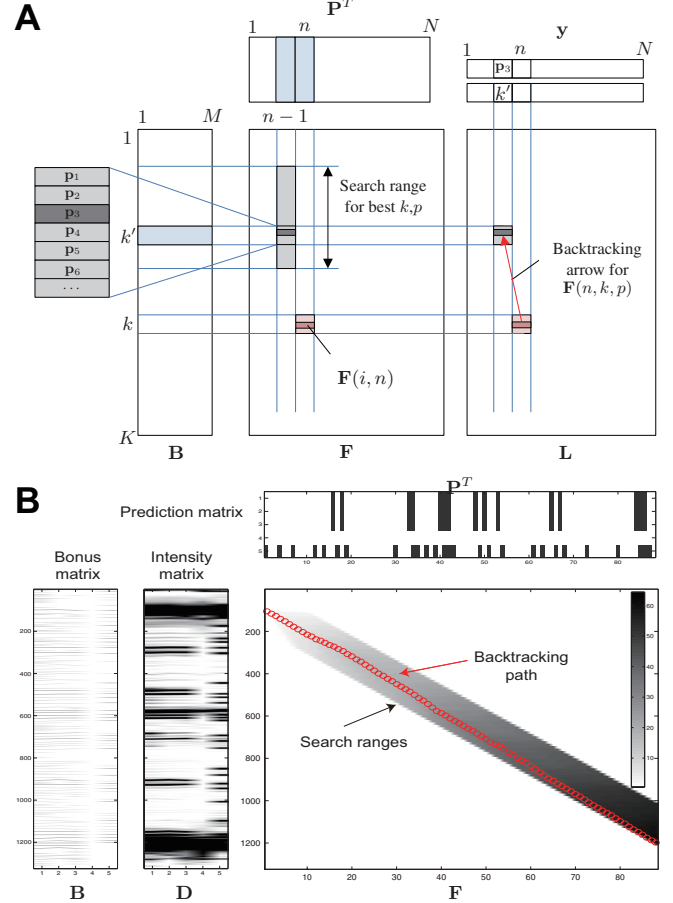
**Fig. 2.** Prediction matrix. **(a)** Definition of the values appearing in the peak prediction matrix. 1 means that a band is expected in that residue position, whereas 0 means that no band is expected. **(b)** Example target sequence and its structure predicted by the Vienna RNA package (Hofacker, 2003). **(c)** The prediction matrix for the example in **(b)**.

exponential increase of time complexity according to the increase of  $M$ . An alternative approach is not to involve peak matching problem in the dynamic programming and assign peak locations to each band only based on the alignment of the other two sequences and heuristic rules. In this paper, a simple rule is proposed where each band is assigned the nearest peak with acceptably short distance. It is a simple and intuitive rule, but there is no way to prevent a single peak from being simultaneously assigned to two consequent bands under this rule. In order to compromise between fully dynamic programming approach and heuristic rule-based approach, a primary lane is selected and its peaks are assigned to bands by the dynamic programming whereas peaks of the other lanes are assigned to bands by the heuristic rule. For notational simplicity, the last lane  $\mathbf{d}_M$  is considered as the primary lane in the rest of this paper.

More formally, define a 3-dimensional matrix  $\mathbf{F}$  indexed by  $n$ ,  $k$  and  $p$  ( $1 \leq n \leq N$ ;  $1 \leq k \leq K$ ;  $-\rho/2 < p < \rho/2$ ) where the value  $\mathbf{F}(n, k, p)$  indicates the maximum score up to the band  $n$ , position  $k$ , and primary peak - peak on the primary lane - at  $k + p$ . The matrix  $\mathbf{F}$  is filled up recursively:

$$\mathbf{F}(n, k, p) = \max_{\substack{k-2.5\rho \leq k' < k \\ |p| < \rho/2 \\ k' + p' < k + p}} \{ \mathbf{F}(n-1, k', p') + S(n, k', k, p) \} \quad (1)$$

where  $S(n, k', k, p)$  is the score gained by going from position  $k'$  to  $k$  for band  $n$  and primary peak at  $k + p$ . The first constraint on  $k'$  in (1) implies that the jump from  $k'$  to  $k$  must be forward and the distance between them is capped by a reasonable upper bound so that the entire search space can be narrowed down into a probable area for efficient implementation. The search space is reduced even further into the trace of a moving window as shown in Figure 3, during our implementation. The constraint  $k' + p' < k + p$  means that the corresponding primary peak must move forward as well as position  $k$ . More details of  $S(n, k', k, p)$  is described in the next section.



**Fig. 3.** Formulation as dynamic programming. **(a)**  $\mathbf{F}(n, k, p)$  depends on  $\mathbf{F}(n-1, k', p')$  in the previous column and the gap bonus  $S(n, k', k, n)$  between them. The best tuple  $(k', p')$  that maximizes  $\mathbf{F}(n, k, p)$  is searched for in the range  $k - 2.5\rho \leq k' < k$ ;  $k' + p' < k + p$  and is stored in the backtracking matrices  $\mathbf{L}_k(n, k, p)$ ,  $\mathbf{L}_p(n, k, p)$ . The computation of  $S(n, k', k, p)$  is based on the bonus matrix  $\mathbf{B}$  and the prediction matrix  $\mathbf{P}$  (Section 2.6). **(b)** Example. The data set used is xxx.  $N = 88$ ,  $M = 5$ ,  $K = 1324$ . The backtracking path is represented by a series of red circles superimposed on the score matrix  $\mathbf{F}$ ; since  $\mathbf{F}$  is 3-dimensional, the figure alternatively represents a reduced matrix  $\mathbf{F}'$  defined by  $\mathbf{F}'(n, k) = \max_{p'} \mathbf{F}(n, k, p')$ . The output array  $\mathbf{y}_k$ , which stores the position of each circle, indicates the band locations.

The backtracking matrices  $\mathbf{L}_k, \mathbf{L}_p$  for finding the solution itself are given by

$$\begin{aligned} \mathbf{L}(n, k, p) &= (\mathbf{L}_k(n, k, p), \mathbf{L}_p(n, k, p)) \\ &= \underset{\substack{k-2.5\rho \leq k' < k \\ |p| < \rho/2 \\ k' + p' < k+p}}{\operatorname{argmax}} \{ \mathbf{F}(n-1, k', p') + S(n, k', k, p) \} \end{aligned} \quad (2)$$

and respectively store the position  $k$  and the relative peak location  $p$  from which  $\mathbf{F}(n, k, p)$  is derived as in (1). The output array  $\mathbf{y}$  is derived from  $\mathbf{L}_k$  and  $\mathbf{L}_p$  as follows:

$$\begin{aligned} \mathbf{y}(n) &= (\mathbf{y}_k(n), \mathbf{y}_p(n)) \\ &= \begin{cases} \operatorname{argmax}_{k,p} \{\mathbf{F}(N, k, p)\}, & \text{if } n = N; \\ \mathbf{L}(n+1, \mathbf{y}_k(n+1), \mathbf{y}_p(n+1)), & 1 \leq n \leq N-1. \end{cases} \end{aligned} \quad (3)$$

The value of  $\mathbf{y}_k(n)$  corresponds to the location of the  $n$ -th band in the input sequence  $\mathbf{s}$ . Figure 3 illustrates the proposed dynamic-programming formulation with an example.

## 2.6 Description of score term

The score term in (1) consists of the following two components:

$$S(n, k', k, p) = S_{\text{dist}}(n, k - k') + w_{\text{peak}} \cdot S_{\text{peak}}(k, p) \cdot \mathbf{P}(n, :) \quad (4)$$

where  $S_{\text{dist}}$  and  $S_{\text{peak}}$  are functions returning nonnegative values and  $\mathbf{P}(n, :)$  is the  $n$ -th row of the prediction matrix  $\mathbf{P}$ .

**2.6.1 Distance bonus term** It is empirically supported that the length between consecutive locations,  $k'$  and  $k$ , is quite evenly distributed.  $S_{\text{dist}}$  is the bonus term that utilizes this fact and induces the dynamic programming to end up with regularly stretched output. In addition, observations on reference annotations suggest that a gap between two consecutive locations tends to be shorter when the preceding location corresponds to 'G' in the RNA sequence. These observations lead to the definition of distance bonus term as follows:

$$S_{\text{dist}}(n, d) = \frac{f_{(\rho', \frac{\rho}{2})}(d)}{f_{(\rho', \frac{\rho}{2})}(0)} \quad (5)$$

where

$$\rho' = \begin{cases} \frac{2}{3}\rho, & \text{if } \mathbf{s}(n-1) = \text{G}; \\ \rho, & \text{otherwise} \end{cases}$$

and  $f_{(\mu, \sigma)}$  is the density function of  $N(\mu, \sigma)$ . That is,  $S_{\text{dist}}(n, d)$  reaches its maximum value 1 when  $d = \rho'$  and decreases along a Gaussian curve as  $d$  deviates from  $\rho'$ .

**2.6.2 Peak bonus term** Another bonus term used in this method is the peak bonus term which is used to incline our output to locate bands near peaks with a significant curvature. As the peak bonus is granted only for the lanes with a band at the location,  $P(n, :)$  must be referred before actually adding up the bonuses as demonstrated in the equation (4).  $S_{\text{peak}}$  is a function that returns a nonnegative  $M$ -dimensional value where each of its entries represents the peak bonus from each lane:

$$S_{\text{peak}} = (S_{\text{peak}}^1, S_{\text{peak}}^2, \dots, S_{\text{peak}}^M) \quad (6)$$

where  $S_{\text{peak}}^m$  stands for the bonus from matching a peak to a band in  $\mathbf{d}_m$ , assuming such a band exists. The bonus is boosted for a greater curvature at the peak and the proximity of the peak to the band, so  $S_{\text{peak}}^m(k, p)$  is defined as the product of a Gaussian density function and an entry of  $\mathbf{B}$  corresponding to the peak:

$$S_{\text{peak}}^m(k, p) = \frac{f_{(0, \frac{\rho}{5})}(\tilde{p})}{f_{(0, \frac{\rho}{5})}(0)} \cdot \mathbf{B}(k + \tilde{p}, m) \cdot \alpha_m \quad (7)$$

where

$$\alpha_m = \begin{cases} M^{\frac{1}{2}}, & m = M; \\ 1, & \text{otherwise} \end{cases}$$

and

$$\tilde{p} = \begin{cases} p, & \text{if } m = M; \\ \underset{q \in [-5, 5]}{\operatorname{argmin}} |q|, & \text{otherwise} \\ \mathbf{B}(k+q) \neq 0 \end{cases}$$

If such  $\tilde{p}$  does not exist, set  $S_{\text{peak}}^m(k, p) = 0$ . As shown above, the peak bonus from the primary lane is given more weight because it plays an important role in finding the optimal solution with dynamic programming.

**2.6.3 Peak bonus weight coefficient** Two distinct bonus terms are normalized by a linear combination, where an optimal selection of weight coefficient is crucial. An overvalued weight on peak bonus is most likely to result in solutions with exact matches of bands to peaks and uneven distribution of gaps whereas an undervalued weight  $w_{\text{peak}}$  would bring about output with uniform distances neglecting peak-band matching. An optimal coefficient  $w_{\text{peak}}$  varies according to dataset and can be discovered through iterative calculations. For the dataset used here,  $w_{\text{peak}} = 1.5$  leads to the best output.

**Table 1.** High-throughput RNA structure mapping data sets analyzed by the proposed method (total 522 profiles and 47210 bands). Excluding the last line, there are 95 data sets. More details of these 95 data sets are described in Lee *et al.* (2012). The last data set is from a study on a 187-nt ribozyme.

Name	# profiles	# nt	# bands per profile	# total bands
R45 <sup>a</sup>	60	108	88	5280
R46 <sup>a</sup>	80	108	88	7040
R47 <sup>b</sup>	90	112	92	8280
R47B <sup>b</sup>	36	112	92	3312
R48 <sup>b</sup>	96	112	92	8832
R49 <sup>b</sup>	18	112	92	1656
R49B <sup>c</sup>	48	115	95	4560
R50 <sup>c</sup>	54	115	95	5130
R43 <sup>d</sup>	40	98	78	3120
HDV <sup>e</sup>	xx	187	yy	zz

<sup>a</sup> Flavin mononucleotide (FMN) aptamer with single binding site (Lee *et al.*, 2012); <sup>b</sup> FMN aptamer with single binding site II; <sup>c</sup> FMN binding branches; <sup>d</sup> The backwards C; <sup>e</sup> NMIA (SHAPE) modification of the hepatitis delta virus (HDV) ribozyme

## 2.7 Reliability evaluation

Regardless of how accurately the automated band annotation works, it may always return a misleading output. Thus, it must be supported by a method to measure the reliability of automatically generated band locations prior to practical application. The focus is on the intuition that when the dynamic programming fails to approach the desirable solution for any reason, the balance between two bonus terms is broken because the value of the weight coefficient  $w_{\text{peak}}$  itself originates from the nature of the data and its correct answer. There are two possible consequences when  $w_{\text{peak}}$  fails to arbitrate between the distance bonus term and the peak bonus term: extraordinarily short or long distances between consecutive locations, or bands on the primary lane without proper matching to peaks. We devise an indicator for the reliability of output based on the number of such occurrences as follows:

- E1: number of bands without corresponding peak
- E2: number of gaps with length less than or equal to  $\rho/4$
- E3: number of gaps with length greater than or equal to  $2\rho$
- ME = max(E1, E2, E3)

Greater ME score means more abnormalities in the output which is believed to result from output digressing from the correct answer, so it can be expected that output with the smaller ME would be more reliable than output with the greater ME. The relationship between the ME score and the actual reliability of output is discussed in the result section.

## 2.8 Implementation and data preparation

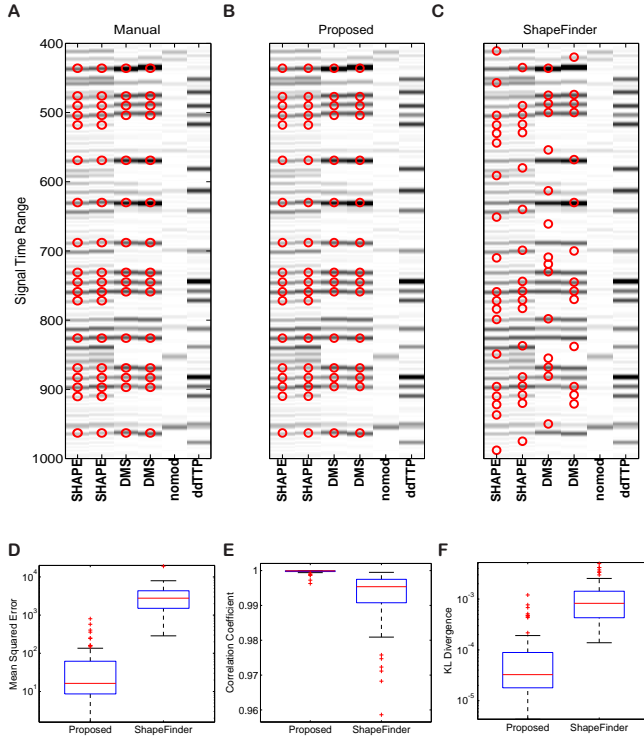
We implemented the proposed method in the MATLAB programming environment (The MathWorks, <http://www.mathworks.com>) and are making it freely available for download at <http://hitrace.stanford.edu>. (To be filled with data preparation...)

## 3 RESULTS

### 3.1 Robust determination of band positions

Figure 4(a)–(c) shows the electrophoretic profiles annotated with band locations by three different methods: reference, proposed,





**Fig. 4.** Determination of band locations. (A) Reference (manual) annotation. Red circles represent band locations. (B) The band locations determined by the proposed method. (C) The band locations found by ShapeFinder (Vasa *et al.*, 2008). (D) Mean squared error (MSE) (E) Pearson's correlation coefficient (F) KL divergence

and ShapeFinder (Vasa *et al.*, 2008), respectively. The reference annotation was prepared manually and verified by experimentalists. Visual inspection suggests that the proposed method produces annotations more compatible with the reference. The annotations determined by ShapeFinder tend to be off from the reference positions downwards. We could observe this trend not only in this profile but in general. In addition, ShapeFinder seems to have difficulties in locating some band locations. For the 95 data sets used in the experiments, ShapeFinder missed  $zz\%$  annotations on average per data set. This limitation originates from the fact that ShapeFinder does not take the number of bands to be annotated as input but determines the band locations based on the intensity and reference ladder. For fair comparison, we considered only those band locations both ShapeFinder and the proposed method identifies, not to penalize ShapeFinder for failing to mark some band locations. Supplementary Material has the band annotation plots for all the profiles used in the experiment.

To assess the accuracy of automated band annotation quantitatively, we computed the mean squared error (MSE) of the band locations determined by the proposed method with respect to the reference locations. In addition, we measured the Pearson's correlation coefficient and the KL divergence between the reference and computationally determined band positions. For comparison, we repeated calculating quantitative metrics for ShapeFinder over the same data. The box plots in Figure 4(d)–(f) reveal that the proposed method outperforms ShapeFinder in all the three

metrics by large margin:  $xx\%$ ,  $yy\%$ , and  $zz\%$  in terms of MSE, correlation coefficient, and KL divergence, respectively. According to our experience, a band annotation result with MSE lower than 100 is practically indistinguishable from the reference. For the proposed method, most MSE values were lower than 100 with the median MSE of  $xx$ . In contrast, nearly all MSE values were above 100 for ShapeFinder (median:  $yy$ ). This suggests that more manual intervention would be needed if ShapeFinder were used as the band annotation tool. The MSE, correlation coefficient, and KL divergence values for individual profiles are provided in Supplementary Material.

### 3.2 Accurate peak-area quantification

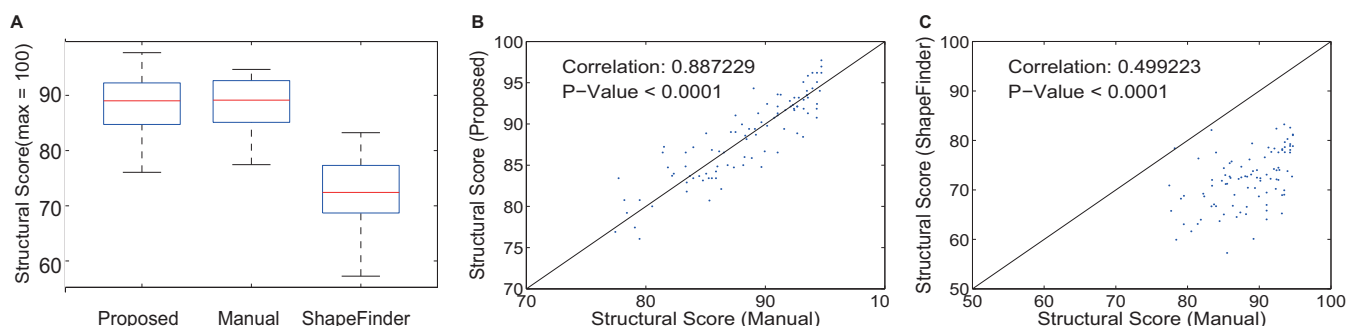
In the RNA structure mapping pipeline, the band annotation is followed by peak deconvolution, which fits each band with a Gaussian curve and produces the quantified area of the band. To see how the final band quantification results get improved by employing the proposed band annotation method, we calculated Pearson's correlation coefficients between band intensities quantified based on the band annotation found by the proposed method and those quantified based on the reference annotation. We also repeated the calculation with the band intensities quantified by ShapeFinder. For fair comparison, we applied the same peak deconvolution software (HiTRACE; Yoon *et al.*, 2011) to these three methods.

Figure 5(a) shows the distribution of the Pearson's correlation coefficients measured using the entire 95 data sets. The median correlation coefficient for the proposed method is  $zz\%$  higher than that for ShapeFinder, and the distribution for the proposed method shows smaller variance. This observation suggests that using the proposed band annotation can significantly enhance the accuracy of band quantification. Figure 5(b) and (c) shows the correlation of results between the proposed method and reference for a specific data set (RRRR), and Figure 5(d) and (e) shows the correlation between the ShapeFinder and reference results. As expected from Figure 5(a), the proposed method produces results that are more correlated with the reference than ShapeFinder. See Supplementary Material for the results from all individual data sets.

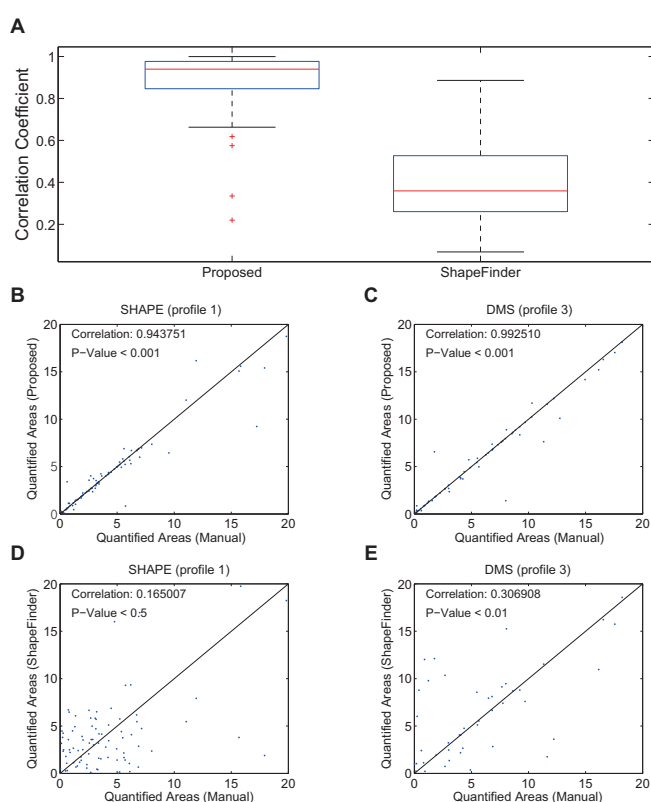
### 3.3 High consistency with predicted structure

We utilize the scoring scheme designed to assess the consistency of quantified bands with predicted structure, as described in (Lee *et al.*, 2012). Briefly, the scoring procedure proceeds as follows: From the quantified band areas in each capillary, we subtract the corresponding areas in the control capillary. We normalize the resulting areas and fit them with the corresponding entries in the band prediction matrix  $\mathbf{P}$  (see Section 2.3) by scaling and baseline adjustment. Note that elements in  $\mathbf{P}$  can have either zero or one, representing a paired and unpaired residue, respectively. For each band, if the prediction is one (zero), then the band scores one if its adjusted intensity is higher (lower) than a threshold. The scores of all bands in a capillary is accumulated and normalized to the number of bands in that capillary, giving a total score between 0 and 100 percent.

In Figure 6(a), we compare the distributions of the scores calculated from the band locations determined by the proposed method, ShapeFinder and the reference. For each method, we quantified hand areas by the same HiTRACE software. Evidently, the distribution of the scores from the proposed method seems



**Fig. 6.** Consistency with predicted structure. We utilized a scoring scheme described in Lee *et al.* (2012) that can assess how closely the band configuration in a CE profile matches its target secondary structure. **(a)** The distributions of structure scores measured over the 95 data sets. To see the performance gap at individual data-set level, **(b)** and **(c)** show the correlation between the reference scoring scores and those by the proposed method and ShapeFinder, respectively.



**Fig. 5.** Accuracy of quantifying peak areas. **(a)** The left box plot represents the distribution of the Pearson's correlation coefficients between the manually quantified areas and those quantified by the proposed method over the 95 data sets. The right box plot shows the distribution of the correlation coefficients between the areas quantified manually and by ShapeFinder. **(b–c)** Correlation of the reference and the quantified areas by the proposed method for data set RRRR is shown for profiles 1 (SHAPE) and 3 (DMS). **(d–e)** Correlation of the reference and the areas quantified by ShapeFinder for the same data set.

highly close to that of the reference scores. In contrast, the score distribution for ShapeFinder results is positioned significantly lower. Given that all three distributions have nearly identical variances,

we carried out a statistical test to compare the means of the three distributions. Based on the t-test, we were able to reject the null hypothesis that the means of the two distributions from the reference and ShapeFinder are identical at 0.001 level. In contrast, we could not reject the null hypothesis for the distributions from the proposed method and the reference at the same level.

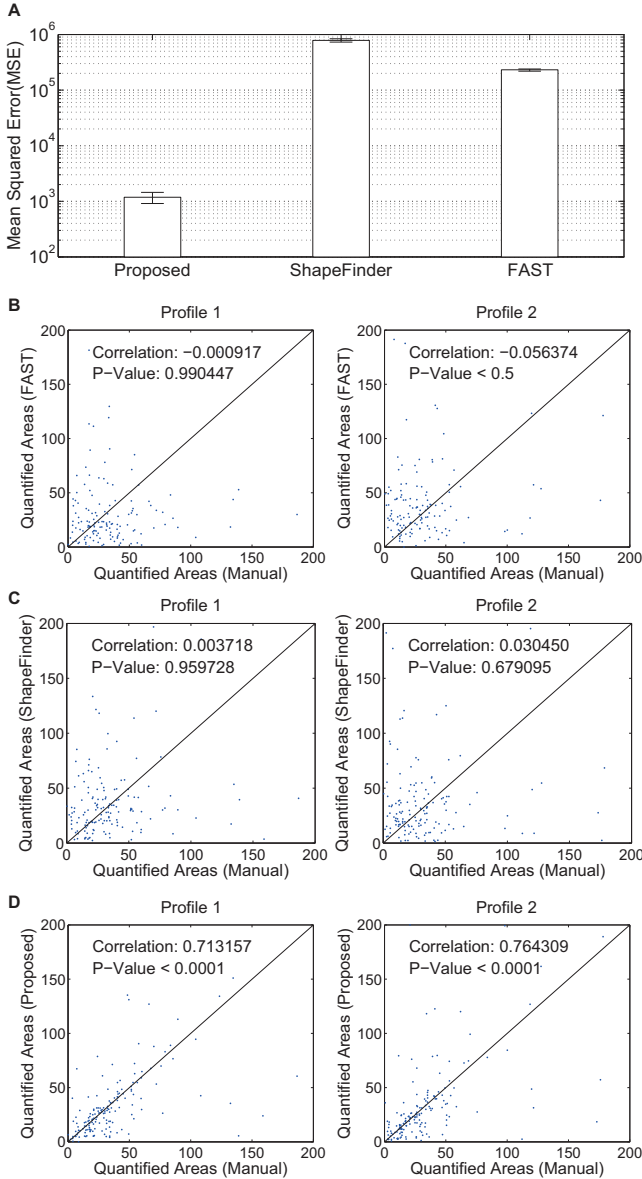
To see the performance difference at individual data-set level, Figure 6(b) and (c) presents the correlation between the reference structure scores and those of the proposed method and ShapeFinder, respectively, for a specific data set (RRR). The proposed method showed a higher correlation coefficient than ShapeFinder (0.887 versus 0.499). We could observe similar performance gaps in the other data sets (see Supplementary Material for details).

### 3.4 Additional results in longer RNAs

To test the proposed method with a longer RNA molecule, we used a chemical mapping data set from a study on the NMIA (SHAPE) modification of the hepatitis delta virus (HDV) ribozyme (187-bp). For this data set, we included the FAST software (Pang *et al.*, 2011) in comparison. FAST requires ddGTP as reference ladder and could not be used for the other 95 data sets presented earlier (they contain ddTTP profiles instead).

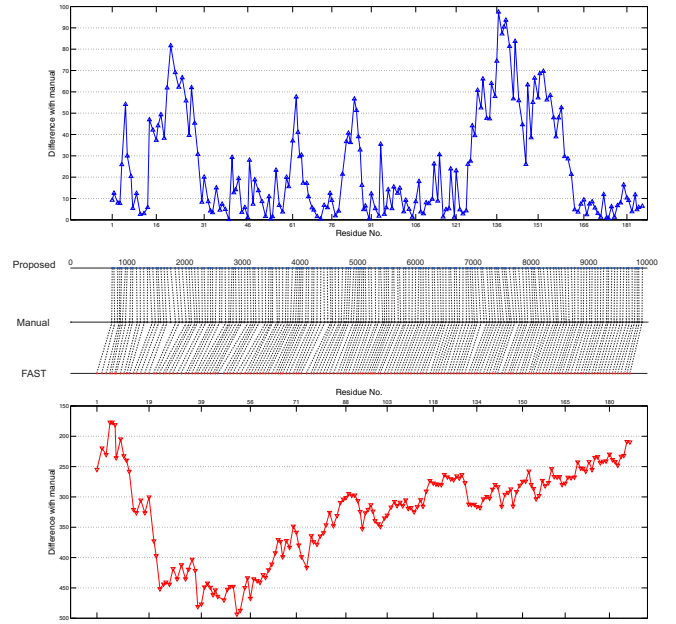
We repeated the experiments presented in Sections 3.1 and 3.2 for this HDV data set. Figure 7(a) shows the the average MSE values of band positions determined by the three methods under comparison. Compared to the reference band locations, the proposed method produced the least amount of error, although its average MSE tends to be higher than those for shorter sequences (Figure 4(d)). Still, the proposed method substantially outperformed FAST and ShapeFinder. Figure 7(c)-(d) shows the correlation of the reference areas with band areas quantified by the proposed method, ShapeFinder, and FAST, respectively. The band areas quantified by the proposed method show a high degree of correlation with the reference ( $r = 0.713$  and  $0.764$ ), whereas the two other methods give unsatisfactory results ( $r < 0.05$ ).

Figure 8 shows the distribution of errors in predicted band locations over the position of residues in the HDV data set. This is to check if there is any systematic bias in band annotation on the residue position. The middle diagram represents the mapping between the reference band locations and those determined by the proposed method. The upper and lower plots show the position



**Fig. 7.** Assessing band annotation results from 187-nt HDV data set. (a) The average MSE between the reference band locations and those determined by different approaches. (b–d) Correlation between the areas quantified manually and by FAST, ShapeFinder, and the proposed method over two profiles, respectively.

errors for the proposed method and FAST, respectively. For the proposed method, the errors near the start and end locations tend to be larger than those in the middle. According to our experience, there exist very high-intensity bands in the starting and ending portion of a profile, which hinders even the manual band annotation. The larger error near these segments in a profile may be due to these high-intensity bands. The error pattern in the result from FAST shows a different pattern, and the largest errors appear near residues 40–50. Overall, the errors from the proposed method were consistently lower than those from FAST.



**Fig. 8.** Error in band positions with respect to the reference band locations for 187-nt HDV data. Upper plot: error over residue positions for the proposed method; middle: mapping between the reference and computationally predicted band locations; lower: error over residue positions for FAST.

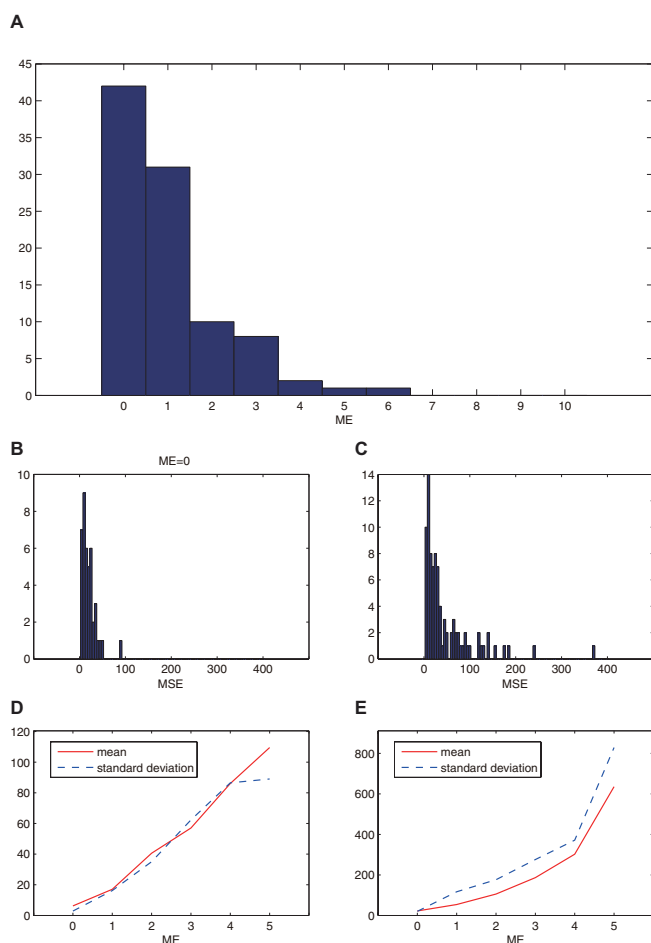
### 3.5 Strong correlation between ME score and MSE

Figure 9(A) shows the histogram of the ME scores over the 95 data sets prepared. Approximately 45 percent of the data sets have zero ME score and about 30 percent of the data sets have ME score one, implying that the proposed method gives us output with fairly low ME scores. Figure 9(B)–(C) show the histograms of the MSE of the output with zero ME scores and the MSE of the whole output respectively. It is obvious from these histograms that MSEs are clustered near 0 when the ME score is zero, allowing us to believe that ME score has a significant influence on the distribution of MSE. Since 95 data sets are not sufficient to conclude on the statistical correlation, however, artificial data sets are generated based on the original data sets through random convolution in terms of amplitude and interval for further verification. Figure 9(D)–(E) show the trend of mean and standard deviation of MSE with respect to ME score, where (D) comes from artificial data generated from a single data set whereas artificial data involved in (E) is generated from the whole 95 data sets. Now that they both show increasing mean and standard deviation of MSE with increasing ME score, it is concluded that the ME score is a strong indicator for the reliability of band locations given by the proposed method.

## 4 DISCUSSION

### 4.1 Use of RNA secondary structure information for band annotation

In the band annotation procedure for an RNA sequence, the proposed method constructs the band prediction matrix  $\mathbf{P}$  based on the RNA's secondary structure predicted by the Vienna RNA



**Fig. 9.** (a) The distribution of ME scores over 95 data sets. (b) The distribution of MSE for the results with zero ME scores. (c) The distribution of MSE for the whole results. (d) The trends of mean and standard deviation of MSE with respect to ME score over artificially generated data sets from a single original data set. (e) The trends of mean and standard deviation of MSE with respect to ME score for artificially generated data sets from the whole 95 data sets.

package (Hofacker, 2003). Although the secondary structure prediction software typically matches most experimental profiles closely, there may exist cases (*e.g.*, complicated pseudoknots) in which the prediction quality is low or even fails. In such cases, using secondary structure information may lead to incorrect band annotation, but through preliminary filtering we can reduce the possibility of inaccurate annotation.

## 4.2 Algorithm complexity and time demand of band annotation

The proposed algorithm relies on dynamic programming and a simple implementation would have the worst-case time and space complexity cubic to the input size. Even so, the practical time demand of band annotation was tolerable in our experiments. Under the experimental setup used (sequential execution on a Intel core i5 4570 processor with 8-GB main memory), the total time demand of

annotating bands in all the 95 data sets did not exceed 5 min (for each data set, mean 2.8517 sec; median 2.8413 sec).

## 4.3 Conclusion

In the analysis of CE profiles, the band annotation is one of the most time-consuming steps, due to the lack of robust computational tools for automated annotation. Using a dynamic-programming approach, the proposed algorithm can find an optimal arrangements of bands in a given CE profile, under the carefully crafted scoring scheme suitable for high-throughput CE experiments. In addition, the proposed reliability index may be consulted while deciding whether or not to adopt the automatically generated annotation. According to our experiments with nearly 100 CE data sets representing approximately 47,000 band locations in total, the proposed method identified the band positions matching the reference positions closely.

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