iRSpot-Pse6NC: Identifying recombination spots in *Saccharomyces cerevisiae* by incorporating hexamer composition into general PseKNC

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**ABSTRACT**

 Meiotic recombination caused by meiotic double-strand DNA breaks. In some regions the frequency of DNA recombination is relatively higher, while in other regions the frequency is lower: the former is usually called “recombination hotspot”, while the latter the “recombination coldspot”. Information of the hot and cold spots may provide important clues for understanding the mechanism of genome revolution. Therefore, it is important to accurately predict these spots. In this study, we rebuilt the benchmark dataset by unifying its samples with a same length (131 bp). Based on such a foundation and using SVM (Support Vector Machine) classifier, a new predictor called “iRSpot-Pse6NC” was developed by incorporating the key hexamer features into the general PseKNC (Pseudo K-tuple Nucleotide Composition) via the binomial distribution approach. It has been observed via rigorous cross-validations that the proposed predictor is superior to its counterparts in overall accuracy, stability, sensitivity and specificity. For the convenience of most experimental scientists, the web-server for iRSpot-Pse6NC has been established at <http://lin-group.cn/server/iRSpot-Pse6NC>, by which users can easily obtain their desired result without the need to go through the detailed mathematical equations involved.

**Key words：**Recombination spot; 5-step rules; Key hexamers; PseKNC; SVM; Webserver

**1. INTRODUCTION**

Meiotic recombination occurs at each generation in diploid organisms, which is caused by meiotic double-strand DNA breaks (DSBs)[1](**Figure 1**). Meiosis can guarantee not only the stability of the chromosome number of species but also a species evolving mechanism to adapt to the environment changes [2]. Recombination can lead to a change in genetic information between homologous chromosomes. Thus, it is one of main driving forces in genome evolution. The frequency of DNA recombination in some regions is relatively higher as referred to recombination hotspots, while in other regions the frequency is lower referred to the recombination coldspots [3-5].



DNA molecules

Double strand break



Strand invasion

No-crossover

Crossover

**Figure 1:** The schematic drawing to show the meiotic recombination pathways in a DNA system.

There have been many in-depth studies of recombination sites [3; 6-9]. Gerton et al. [3] mapped double-strand break sites on chromosomes in the *Saccharomyces cerevisiae* (*S. cerevisiae*), and found that hotspots were non-randomly associated with regions of high GC base composition, while coldspots were non-randomly associated with the centromeres and telomeres. Some hotspots that require transcription factor binding are called *α* hotspots, and others are called *β* hotspots [3]. Recently, there have been new developments on the research of recombination sites. ChIP experiments showed that substantial Spo11 persists at Rec8 binding sites during DSB formation [10]; PRDM9, as a catalytic H3K4 trimethylated histone trimethylase, is involved in the initiation of recombination and recombination with recombination hot spots [11], found that the regions with high nucleosome occupancy have high recombination rate in the yeast genome [12].

The correct identification of recombination spots can provide important clues for understanding the evolution mechanism. Generally, biochemical experiments can produce accurate information for determine recombination spots. However, with the development of high-throughput sequencing technique, more and more genome data were generated, thus, determining recombination spots with these wet-experiments requires more and more expensive experimental materials and long experimental period. Machine learning-based methods are a good choice for timely and accurately identifying the recombination spots. Up to now, some methods have been developed to identify recombination spot. Jiang *et al*. firstly developed a new model based on gapped dinucleotide composition and random forest (RF) to predict meiotic recombination hotspots and coldspots in *S. cerevisiae* [13]. In the meantime, Zhou *et al*. established an SVM-based model to discriminate hotspots from coldspots in *S. cerevisiae* by using codon composition [14]. Subsequently, Liu *et al*. proposed to use the increment of diversity combined with quadratic discriminant for predicting the recombination spots [15]. Chen *et al*. developed a new DNA sample descriptor called pseudo dinucleotide composition (PseDNC) to improve prediction accuracy for the recombination hotspots and coldspots [16]. According to the concept of PseDNC, Li *et al*. [17] and Qiu *et al*. [18] also developed different prediction models to address this problem. Liu *et al*. incorporated the weight of features into recombination hotspots prediction model [19]. A predictor called iRSpot-DACC was also presented to predict recombination hotspots and coldspots [20]. Recently, the same problem was further investigated by including the Z curve approach [21], and the ensemble learning approach [22].

Although the aforementioned methods could achieve quite encouraging results, further studies are needed due to the following reasons. (i) The DNA samples used to train the models are with different length, which prevents them from establishing a widely useful model because users do not know how long the working length should be used for a query DNA sequence. For example, in using the aforementioned methods to scan a chromosome, we do not know the optimal width of the scan window [23] for the biological sequence concerned. In fact, for the published webserver based on those methods, only a prediction will be given even for a chromosome with a length of thousands base pairs. However, there are many recombination points in the genome. Therefore, most of those models are quite limited for practical applications. (ii) Some works [13; 14; 21; 24] used codon composition or coding region information to formulate DNA samples. However, combination spots are not always located in coding regions. Some non-coding regions may also contain combination spots. Thus, these methods could not identify combination spots in the intergenic regions. (iii) The prediction results are still far from satisfactory yet; the accuracy should be further improved. (iv) Only three webservers were published. For the convenience of most experimental scientists, more user-friendly webservers in this regard are needed.

The present study was devoted to develop a more powerful predictor in this area by considering the aforementioned four issues. To make the new predictor more clear in logical development and more useful in practical application, the Chou’s 5-step rules [25] were followed as reported in a series of recent studies (see, e.g., [26-35]).

**2. MATERIALS AND METHODS**

* 1. **Benchmark dataset: hot/cold spots DNA sequences**

According to the Chou’s 5-step rules, the first prerequisite to establish an effective predictor for a biological system is to construct or select a high quality benchmark dataset. In this study, the raw data was derived from Gerton et al. [3], who used DNA microarray as the single-gene resolution method to estimate the DSBs formation adjacent to each ORF for the *S. cerevisiae loci*. They measured the ratio of DSB-rich probes hybridized to total genomic probes. Based on the experimental data, Jiang et al. [13] constructed a benchmark dataset including 490 recombination hotspots and 591 coldspots.

So far most of the existing models [13-20] were built up based on such benchmark dataset. The length distribution of original samples was shown in **Figure 2**. It was noticed that the length distributed in a wide range from the shortest one of 131 bp to the longest one of thousands bp. To overcome such a shortcoming, we rebuilt the benchmark dataset according to the strategy that recombination hotspots were correlated with peaks of G+C base composition [3]. By doing so, we unified the length of each sample to 131 bp because the length of shortest sequence is 131 bp. For those sequences with >131 bp, we chose their subsequences with 131 bp that have the maximum GC content. As a result, the new dataset also has 490 samples for recombination hotspots and 591 samples for recombination coldspots, but all the sequences are 131 bp long now. The new benchmark dataset can be downloaded from the link at <http://lin-group.cn/server/iRSpot-Pse6NC>.

**Figure 2:** The length distribution of benchmark dataset samples.

* 1. **Hexamer composition and its PseKNC vector**

 How to translate a DNA sequence ***D*** with *L* bases into a vector is the second important step to develop a predictor for discriminating recombination hotspots from recombination coldspots. This is because all the existing machine-learning algorithms can only handle vectors but not sequences as elaborated in [36]. But a vector in a discrete framework might totally lose all the sequence-order or pattern information. To deal with this problem, the PseAAC (Pseudo Amino Acid Composition) was introduced [37]. Ever since the concept of PseAAC was proposed, it has been swiftly penetrated into many biomedicine and drug development areas [38; 39] as well as nearly all the areas of computational proteomics (see, e.g., [40-48] and a long list of references cited in a recent review paper [49]). Encouraged by the successes of using PseAAC to deal with protein/peptide sequences, its idea has been extended to deal with DNA/RNA sequences [16; 22; 24; 32; 50] in computational genomics via PseKNC (Pseudo K-tuple Nucleotide Composition) [51; 52]. According to [53], for a DNA sample with *L* nucleic acid residues

$D= R\_{1}R\_{2}R\_{3}\cdots R\_{i}\cdots R\_{L}$ (1)

its general form of PseKNC can be formulated as

$D=\left[ϕ\_{1} ϕ\_{2} \cdots ϕ\_{u} \cdots ϕ\_{Γ}\right]^{T}$ (2)

where **T** is the transposing operator, the subscript $Γ$ is an integer, and its value and the components $ϕ\_{u} \left(u=1, 2, \cdots \right)$ will depend on how to extract the desired features and properties from the DNA sequence. In this study, their definitions are described below.

 K-tuple (or called K-mer) nucleotide composition has important biological significance [54] that the whole DNA sequence can be uniquely determined from the K-tuple nucleotide frequency distribution; i.e., the frequency distribution of K-tuple nucleotide contains mostly the information of the DNA sequence. And K-mer nucleotide composition has been widely used in gene identification [55] and other regulatory element recognition [24; 56-59]. Several studies [60,61] have shown that hexamer (6-mer) distribution has unique properties among species and different DNA fragments. Thus, we have the dimension of PseKNC in Eq.2 is

$Γ=4^{K}=4^{6}=4096$ (3)

and its components given by

$ϕ\_{u}=\frac{n\_{u}}{\sum\_{i=1}^{4096}n\_{i}}=\frac{n\_{u}}{(L-K+1)}$ (4)

where $u$ and *L* denote the number of the *u-*th hexamer and the length of the sample sequence, respectively. Thus, the DNA sample has been uniquely defined in a 4096-D PseKNC vector.

* 1. **The rule for ranking features**

The DNA sequence is represented by a set of 4096 features, which may bring out three problems [62-63]: (1) containing some redundant or irrelevant information; (ii) leading to an over-fitting model and reducing its flexibility; (iii) causing the curse of dimensionality and dyscalculia. However, we can improve these problems by means of the feature selection approach [64]. Many effective feature selection techniques have been proposed, such as diffusion Maps [65], principal component analysis (PCA) [66-68], analysis of variance (ANOVA) [69; 70], recursive feature elimination algorithm [71; 72] and geometry preserving projections (GPP) [73] and so on. These techniques are all quite efficient in alleviating the interference from noise or irrelevant features so as to improve the prediction quality.

Here, let us define a prior probability given by

$P\_{i}=\frac{m\_{i}}{M}$ $(i=1 or 2)$ (5)

where *M* is the total occurrence times of all hexamers in the benchmark dataset (including both positive and negative samples), and $m\_{i} $represents the number of hexamers in the *i*-th type with *i* = 1 referring to the positive subset whereas *i*=2 referring to the negative subset.

Now, the probability of the *j*-th hexamers occurring in type *i* can be formulated as

$P(n\_{ij})=\sum\_{m=n\_{ij}}^{N\_{j}}\frac{N\_{j}!}{m!\left(N\_{j}-m\right)!}P\_{i}^{m}(1-P\_{i})^{N\_{j}-m}$ (6)

where $N\_{j}$ represents the total occurrence number of a given *j*-th hexamer in the benchmark dataset. The smaller the *P(*$n\_{ij}$), the lower the probability of the *j*-th hexamer randomly occurring in type *i*, meaning the hexamer has more biological significance. The confidence level (*CL*) of the *j*-th hexamer occurring in *i*-th type of sample is defined by

$CL\_{ij}=1-P(n\_{ij})$ $(i=1 or 2)$ (7)

Suppose

$CL\_{j}=max⁡(CL\_{1,j}, CL\_{2,j})$ $(j=1,2, \cdots , 4096)$ (8)

thus the 4096 hexamers can be ranked according to the values of Eq.8.

**2.4 Support vector machine**

Support vector machine (SVM) is a supervised machine learning algorithm based on statistical learning theory, and has been successfully applied in the field of bioinformatics [74]. The basic idea of SVM is to transform the data into a high dimensional feature space and then determine the optimal separating hyper plane. For a brief formulation of SVM and how it is working, see the papers [75; 76]; for more details about SVM, see a monograph [77]. In this study, we used the free software LIBSVM 3.20, which was developed by Chang and Lin [78]. Due to its good performance for classification, the radial basis kernel function was used to obtain the best classification hyper plane. The two parameters, *C* and *γ*, which were preliminarily optimized through a grid search strategy.

The proposed predictor thus built up is called iRSpot-Pse6NC, where “i” stands for “identify”, “RSpot” for “Recombination Spots”, and “Pse6NC” for “Pseudo 6-tuple Nucleotide Composition”.

**3. RESULTS AND DISCUSSION**

**3.1 Cross-validation**

 To evaluate the quality of a new predictor, one needs to consider the following two things: (i) what metrics should be used to measure its performance? (ii) what test method should be adopted to calculate these metrics? In literature, the following four metrics are usually used to measure a predictor’s quality [79]: (i) overall accuracy (Acc); (ii) stability (MCC); (iii) sensitivity (Sn); and (4) specificity (Sp). But their conventional expressions directly taken from math books are lack of intuition and difficult to understand by most biological scientists. Fortunately, by means of the symbols introduced by Chou in studying signal peptides [23], the four conventional metrics can be converted to a set of intuitive ones [16; 80; 81] as given below:

$\left\{\begin{array}{c} \\\begin{matrix}Sn=1-\frac{N\_{-}^{+}}{N^{+}} 0 \leq Sn\leq 1 \\\begin{matrix}\begin{matrix}Sp=1-\frac{N\_{+}^{-}}{N^{-}}& 0\leq Sp\leq 1 \end{matrix} & \end{matrix}\end{matrix}\\\begin{matrix}\begin{matrix}\begin{matrix}Acc= Λ=1-\frac{N\_{-}^{+}+N\_{+}^{-}}{N^{+}+N^{-}}& 0\leq Acc\leq 1\end{matrix}\end{matrix} \\\begin{matrix}MCC= \frac{1-\left(\frac{N\_{-}^{+}}{N^{+}}+\frac{N\_{+}^{-}}{N^{-}}\right)}{\sqrt{\left(1+\frac{N\_{+}^{-}-N\_{-}^{+}}{N^{+}}\right) \left(1+\frac{N\_{-}^{+}-N\_{+}^{-}}{N^{-}}\right)}} &-1\leq MCC\leq 1 \end{matrix}\end{matrix}\end{array}\right.$ (9)

where $N^{+}$ represents the total number of positive samples **investigated**, while $N\_{-}^{+}$ is the number of positive samplesincorrectly predicted to be of negative one; $N^{-}$ the total number of negative samples investigated, while $N\_{+}^{-}$ the number of the negative samples incorrectly predicted to be of positive one.

As pointed out by many recent publications (see, e.g., [22; 32; 33; 50; 82-90]), the meanings of Sn, Sp, Acc, and MCC have become crystal clear when using Eq.9.

With a set of intuitive metrics, the next thing is how to test their values. As is well known, the independent dataset test, subsampling (or K-fold cross-validation) test, and jackknife test are the three cross-validation methods widely used for testing a prediction method [91]. To reduce the computational cost, in this study we adopted the 5-fold cross-validation (namely K=5), as done by many investigators with SVM as the prediction engine (see, e.g., [24; 26; 92-95]).

**3.2 Comparison with existing methods**

Listed in **Table 1** are the metrics rates (Eq.9) achieved by iRSpot-Pse6NC via the 5-fold cross-validation on the benchmark dataset (cf. Supporting Information S1). For facilitating comparison, listed there are also the corresponding rates obtained by iRSpot-PseDNC [16], iRSpot-KNCPseAAC [18], and IDQD [15] using exactly the same cross-validation method and same benchmark dataset. As we can see from the table, the rates achieved by iRSpot-Pse6NC are remarkably higher than its cohorts in all the four metrics, clearly indicating the proposed predictor is indeed superior to the existing predictors in this area.

**Table 1.** A comparison of the proposed predictor with the existing ones.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Method | Sna | Spa | Acca | MCCa |
| iRSot-Pse6NCb  | 0.7571 | 0.9103 | 0.8408 | 0.6805 |
| iRSpot-PseDNCc | 0.6234 | 0.9052 | 0.7792 | 0.5585 |
| iRSpot-KNCPseAACd | 0.6102 | 0.8951 | 0.7660 | 0.5334 |
| IDQDe | 0.6959 | 0.7509 | 0.7259 | 0.4469 |

aSee Eq.9 for the metrics definition. bProposed in this paper. cFrom [16]. dFrom [18]. eFrom [15]



**Figure 3** The 5-fold cross-validated IFS curve for predicting recombination hotspots and coldspots. An IFS peak of 84.08% was observed when using the top 381 hexamers to perform prediction.

**3.3 Feature analysis**

As mentioned in section 2.3, the dimension for the hexamer vector is 4096, which is too large to avoid the high-dimension problems. To exclude the noise and redundant features, we used the incremental feature selection (IFS) to find out the best feature subset to maximize accuracy. We initially ranked the 4096 hexamers according to Eqs.5-8. Subsequently, the 4096 feature subsets were obtained, in which the first feature subset contained the first hexamer, the second feature subset was produced by adding the second hexamer into the first feature subset, and so on. Thirdly, the SVM with 5-fold cross-validation was adopted to examine the accuracies of 4096 feature subsets. By using Acc as vertical coordinates and feature number as horizontal coordinates, we plotted IFS curve in **Figure 3**. One may notice that the peak of the curve is 84.08%, which is located at horizontal coordinate of 381. This result (84.08%) is dramatically higher than that (71.04%) of all features. Meanwhile, we also dramatically reduced the considered features from 4096 to 381, indicating that our proposed feature selection technique could pick out the optimal hexamers so as to further improve the prediction quality. Accordingly, the 381 hexamers were selected to form the optimal feature subset to train the prediction model.

To further investigate the performance of the optimal model across the entire range of SVM decision values, we drew the ROC curve [96] in **Figure 4**. It shows that the AUC (the Area Under ROC Curve) reaches the value of 0.9084, indicating that the proposed method is quite promising and holds very high potential to become a useful high-throughput tool for predicting recombination spots.



**Figure 4** The ROC curve for identifying recombination spots by using 381 optimal hexamers. The AUC of 0.9084 was obtained in 5-fold cross-validation. The diagonal dot line denotes a random guess with the AUC of 0.5.

For further analyzing the contributions of different features in the prediction model, a heat map [97] was provided (**Figure 5**), which is a graphical representation of a matrix by using different colors according to its *CL* values scaled between 0 and 1. As we can see from **Figure 5**, for the 4096 different hexamers, the majority of them are blue or green, indicating that most of them are irrelevant to the recombination spot recognition.

It can be seen from **Figure 5** thatthose regions with high GC content, e.g., the hexamers CGCCGG, AGCCGG and GCAGCT, GCCGGA, AGTGGG are with the *CL* values ranking top five among all the features and with the confidence level of *CL* > 98.3%.



**Figure 5:** A heat map to illustrate the *CL* of the 4096 different hexamers. The color scale is ranged from blue (low *CL*) through green and yellow to red (high *CL*). See the main text for further explanation. A higher resolution version can be found at <http://lin-group.cn/server/iRSpot-Pse6NC/heatmap2.jpg>.

Moreover, we performed a detail analysis on the 381 optimal hexamers with *CL*>98.3% to investigate the relationship between the features and GC content (**Figure 6**). In this figure, abscissa coordinate denotes the GC content distribution from 0% -100%, and the vertical axis indicates that the percentage of positive and negative samples at the GC content shown on the abscissa. It can be seen from the figure that the optimal hexamers with high GC content have a higher proportion in positive samples, whereas hexamers with lower GC contents have a higher proportion of negative samples. This means that there is a close relationship between GC content and the hot spots, once again proofing that the way we handled the data is fully valid.

**Figure 6:** The graph to show the relationship between the important features and GC content.

**3.4. Web-server and user guide**

As pointed out in [25] and demonstrated in many follow-up publications (see, e.g., [28; 30; 32; 35; 81; 98-116]), user-friendly and publicly accessible web-servers represent the future direction for developing practically more useful predictors. Actually, a new prediction method with the availability of a user-friendly web-server would significantly enhance its impacts [36; 49]. In view of this, the web-server for iRSpot-Pse6NC has been established. Furthermore, to maximize the convenience of most experimental scientists, the step-by-step instructions are given below.

**Step 1**. Open the web server at http://lin-group.cn/server/iRSpot-Pse6NC and you will see the top page of`iRSpot-Pse6NC shown on your computer screen (**Figure 7**).

**Step 2**. Click on the WEB SERVER button to start the prediction. Either type or copy/paste the query DNA sequences into the input box at the center of **Figure 7**. The input sequences should be in the FASTA format. And click on the Submit button to see the predicted result.

**Step 3**. Click on the DOWNLOAD button to download the benchmark data sets used to train and test the iRSpot-Pse6NC predictor.

**Step 4**. Click on the CITATION button to find the relevant papers that document the detailed development and algorithm of iRSpot-Pse6NC.

**Step 5.** Click on the HELP button to view the relevant instructions and the caveat when using it.



**Figure 7.** A semi-screenshot for the top page of the iRSpot-Pse6NC webserver at <http://lin-group.cn/server/iRSpot-Pse6NC>.

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**Competing Interests**

The authors declare that there are no competing interests regarding the publication of this paper.

**Author Contributions**

H.L. conceived and designed the experiments; H.Y., W.R.Q., G.L., F.B.G. and W.C. analyzed the data and implemented SVM. H.Y., H.L. and W.C. established the web-server; H.Y., W.C., K.C.C. and H.L performed the analysis and wrote the paper. All authors read and approved the final manuscript.

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