# Recent advances in fast algorithms for optimal changepoint detection

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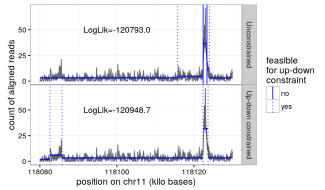
## Constrained Dynamic Programming and Supervised Penalty Learning Algorithms for Peak Detection in Genomic Data

Linear Time Dynamic Programming for Computing Breakpoints in the Regularization Path of Models Selected From a Finite Set

Generalized Functional Pruning Optimal Partitioning (GFPOP) for Constrained Changepoint Detection in Genomic Data

#### Citation

Hocking TD, Rigaill G, Fearnhead P, Bourque G. Constrained Dynamic Programming and Supervised Penalty Learning Algorithms for Peak Detection in Genomic Data. Journal of Machine Learning Research 21(87):1-40, 2020.



Main new idea: an optimal algorithm for computing the best changepoints which alternate up and down.



#### Comparison to previous work and novelty

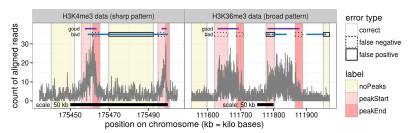
Constraint	No pruning	Functional pruning	
None	Dynamic Prog. Algo. (DPA)	Pruned DPA (PDPA)	
	Optimal, $O(Sn^2)$ time	Optimal, $O(Sn \log n)$ time	
	Auger and Lawrence (1989)	Rigaill (2010); Johnson (2011)	
Up-down	Constrained DPA (CDPA)	Generalized Pruned DPA (GPDPA)	
	Sub-optimal, $O(Sn^2)$ time	Optimal, $O(Sn \log n)$ time	
	Hocking et al. (2015)	This paper	

$$\begin{array}{ll} \underset{0=t_0 < t_1 < \cdots < t_{S-1} < t_S = n}{\text{minimize}} & \sum_{s=1}^{S} \sum_{i=1+t_{s-1}}^{t_s} \ell(u_s, z_i) \\ \text{subject to} & u_{s-1} \leq u_s \ \forall s \in \{2, 4, \dots\}, \\ & u_{s-1} \geq u_s \ \forall s \in \{3, 5, \dots\}. \end{array}$$

- ▶ One hyper-parameter = number of segments  $S \in \{1, 3, ...\}$ .
- ▶ Hard optimization problem, naively  $O(n^S)$  time.
- Previous unconstrained model: not always up-down changes.
- ▶ Interpretable: P = (S 1)/2 peaks (segments 2, 4, ...).
- ▶ H et al., ICML 2015:  $O(Sn^2)$  time approximate algorithm.
- ► This paper:  $O(Sn \log n)$  time optimal algorithm.

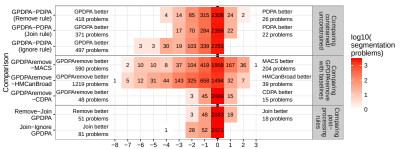


#### Labeled data setting



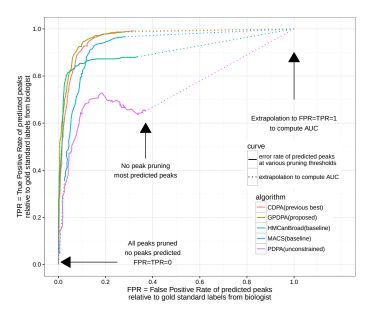
Benchmark data with 2752 labeled data sequences: https://rcdata.nau.edu/genomic-ml/chip-seq-chunk-db/

#### Comparing minimum label errors

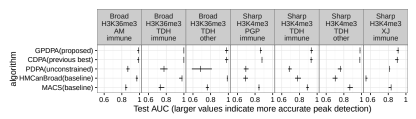


Difference in minimum number of incorrect labels per segmentation problem

#### Test ROC curves of predicted peaks

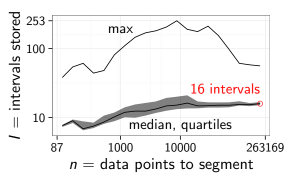


#### Test AUC of predicted peaks

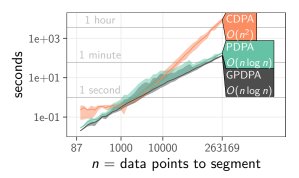


- ▶ 4-fold cross-validation: train on 3/4 of labels, test on 1/4.
- All models trained by learning a scalar significance threshold / penalty parameter, which is varied to compute ROC/AUC.
- MACS is highly inaccurate in all data sets.
- ▶ HMCanBroad is accurate for broad but not sharp pattern.
- Unconstrained PDPA algorithm not as accurate as up-down constrained algorithms (CDPA, GPDPA).

#### Time complexity analysis 1



#### Time complexity analysis 2



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#### Previous work on the Segment Neighborhood problem

Constraint	No pruning	Functional pruning	
None	Dynamic Prog. Algo. (DPA)	Pruned DPA (PDPA)	
	Optimal, $O(SN^2)$ time	Optimal, $O(SN \log N)$ time	
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	Hocking et al. (2015)	This paper	

▶ All algorithms solve the **Segment Neighborhood** "constrained" problem: most likely mean  $m_i$  for data  $z_i$ , subject to the constraint of S segments (S-1) changes.

$$\begin{array}{ll} \underset{\mathbf{m} \in \mathbb{R}^N}{\text{minimize}} & \sum_{i=1}^N \ell(m_i, z_i) \\ \\ \text{subject to} & \sum_{i=1}^{N-1} I[m_i \neq m_{i+1}] = S-1, \end{array}$$

...up-down constraints on m.

#### Previous work on the Optimal Partitioning problem

	no pruning	functional pruning
unconstrained	Opt. Part. Algo.	FPOP
	exact $O(N^2)$	exact $O(N \log N)$
	Jackson et al 2005	Maidstone et al 2016
up-down constrained		Generalized FPOP
		exact $O(N \log N)$
		This work

All algorithms solve the **Optimal Partitioning** "penalized" problem: most likely mean  $m_i$  for data  $z_i$ , penalized by a non-negative penalty  $\lambda \in \mathbb{R}_+$  for each change:

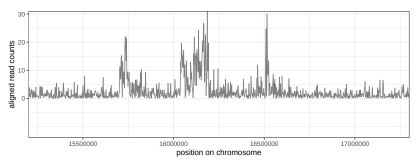
$$\underset{\mathbf{m} \in \mathbb{R}^N}{\text{minimize}} \quad \sum_{i=1}^N \ell(m_i, z_i) + \lambda \sum_{i=1}^{N-1} I[m_i \neq m_{i+1}]$$

subject to ...up-down constraints on m.

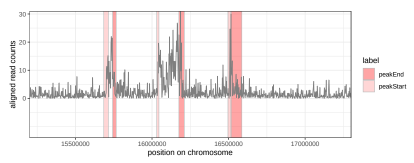
#### Benchmark of large genomic data sequences

http://github.com/tdhock/feature-learning-benchmark

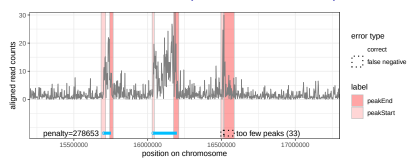
- ▶ 4951 data sequences ranging from  $N = 10^3$  to  $N = 10^7$ .
- ▶ Ran GFPOP with penalty  $\lambda \in (\log N, N)$ , resulting in a range of models with different numbers of peaks, for each data set.
- ► Each data set has **labels** which can be used to determine an appropriate number of peaks.



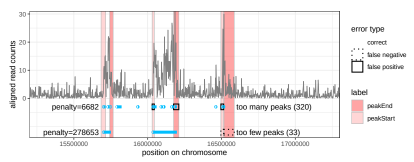
One ChIP-seq data set with N = 1,254,751 (only 82,233 shown).



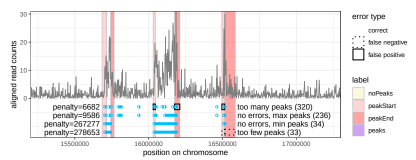
Visually labeled regions (H et al., Bioinformatics 2017).



Penalty too large, too few peaks, 2 false negative labels.

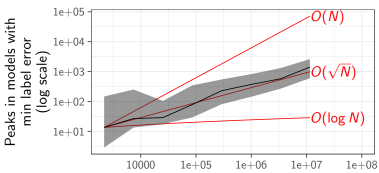


Penalty too small, too many peaks, 3 false positive labels.



Models with 34-236 peaks have no label errors (midpoint=135).

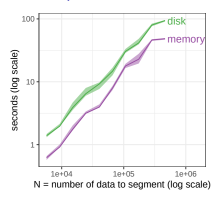
### Segment Neighborhood model too slow for $O(\sqrt{N})$ peaks



N = number of data to segment (log scale)

- Previous GPDPA: O(S) dynamic programming iterations, each is  $O(N \log N)$  time/space.
- If we want  $S = O(\sqrt{N})$  segments then the algorithm is  $O(N\sqrt{N}\log N)$  time/space too much for large data.
- For example  $N=10^7$  data. Each  $O(N \log N)$  DP iteration takes 1 hour, 80 GB. Overall if we want  $S=O(\sqrt{N})=2828$  segments we need means 220 TB of storage space and 17 weeks of computation time!

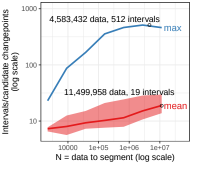
#### GFPOP Implementation stores cost functions on disk

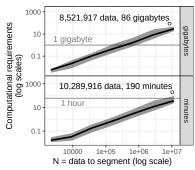


- Disk storage is only a constant factor slower than memory!
- ightharpoonup Both are  $O(N \log N)$  time.
- ▶ Memory implementation:  $O(N \log N)$  memory! (too big)
- ▶ Disk implementation:  $O(\log N)$  memory (< 1GB),  $O(N \log N)$  disk.

### Time/space to solve one penalty is $O(N \log N)$

Total time/space = O(NI) where I is the number of intervals (candidate changepoints) stored in every optimal cost function  $\overline{C}_t(\mu)$ .





 $I = O(\log N)$  intervals.

Overall  $O(N \log N)$  complexity.

But how to compute model with  $O(\sqrt{N})$  peaks?



# Binary search algorithm using GFPOP to compute most likely model with at most $P^*$ peaks

- 1: Input: data  $\mathbf{z} \in \mathbb{R}^N$ , target peaks  $P^*$ .
- 2:  $\overline{L}, \overline{p} \leftarrow \mathsf{GFPOP}(\mathbf{z}, \lambda = 0) // \mathsf{max} \mathsf{ peak} \mathsf{ model}$
- 3:  $\underline{L}, \underline{p} \leftarrow \mathsf{GFPOP}(\mathbf{z}, \lambda = \infty) \ // \ \mathsf{0} \ \mathsf{peak} \ \mathsf{model}$
- 4: While  $p \neq P^*$  and  $\overline{p} \neq P^*$ :
- 5:  $\lambda = (\overline{L} \underline{L})/(p \overline{p})$
- 6:  $L_{\text{new}}, p_{\text{new}} \leftarrow \overline{\mathsf{GFPOP}}(\mathbf{z}, \lambda)$
- 7: If  $p_{\text{new}} \in \{\underline{p}, \overline{p}\}$ : return model with  $\underline{p}$  peaks.
- 8: If  $p_{\text{new}} < P^*$ :  $\underline{p} \leftarrow p_{\text{new}}$
- 9: Else:  $\overline{p} \leftarrow p_{\text{new}}$
- 10: If  $\underline{p} = P^*$ : return model with  $\underline{p}$  peaks.
- 11: Else: return model with  $\overline{p}$  peaks.

### Example run of binary search algorithm using GFPOP

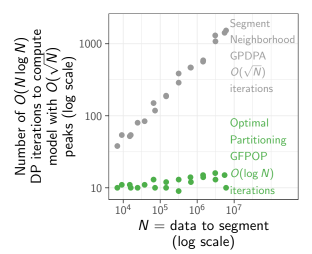
One data set with  $P^* = 93$  peaks and N = 146, 186 data.

iteration	<u>p</u>	$\overline{p}$	$\lambda$	$p_{new}$	$L_{new}$
1			0	68752	-2570319
1			$\infty$	0	14239212
2	0	68752	244.4952	4361	1980119
3	0	4361	2811.0738	188	3676671
4	0	188	56183.7271	55	5330310
5	55	188	12433.3766	98	4108354
6	55	98	28417.5941	72	4584042
7	72	98	18295.6895	83	4336773
8	83	98	15227.9249	90	4218815
9	90	98	13807.6282	95	4146172
10	90	95	14528.5052	92	4188881
11	92	95	14236.0863	94	4160179
12	92	94	14350.6622	93	4174480

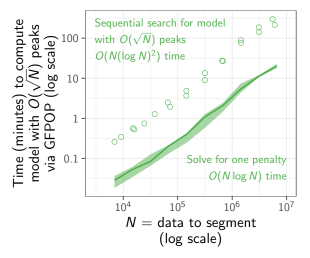
12 DP iterations much fewer than  $93 \times 2 = 186$  which would be required for Segment Neighborhood!



### Only $O(\log N)$ runs of GFPOP to compute $O(\sqrt{N})$ peaks



# Binary search only a log factor slower than solving one penalty



For  $N = 10^7$  only several hours of computation! (compare with weeks for Segment Neighborhood algorithm)

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#### Conclusions

- Previous GPDPA was  $O(N\sqrt{N}\log N)$  much too complex to compute a zero-error model with  $O(\sqrt{N})$  peaks for  $N=10^7$  data.
- ▶ Proposed GFPOP with binary search is  $O(N(\log N)^2)$  time,  $O(\log N)$  memory,  $O(N \log N)$  disk optimal models are now possible to compute for large data.
- C++ code with R interface: PeakSegPipeline::PeakSegFPOP\_disk https://github.com/tdhock/PeakSegPipeline
- Future work: changepoint detection in large ecological data?
- Contact me: toby.hocking@nau.edu
- ▶ Thanks!

### For some data the desired number of peaks does not exist!

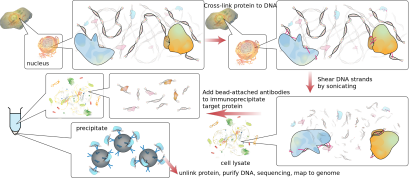
One data set with  $P^* = 75$  and N = 66,031 data.

itera	ation	<u>p</u>	$\overline{p}$	$\lambda$	$p_{new}$	$L_{new}$
	1			0	29681	-1495863.85
	1			$\infty$	0	1200631.42
	2	0	29681	90.85	3445	-1245181.37
	3	0	3445	709.96	401	-446632.57
	4	0	401	4107.89	51	3105.03
	5	51	401	1284.96	168	-230152.31
	6	51	168	1993.65	97	-120725.83
	7	51	97	2691.98	68	-53946.18
	8	68	97	2302.75	81	-86423.80
	9	68	81	2498.28	77	-76891.10
	10	68	77	2549.44	71	-61722.09
	11	71	77	2528.17	74	-69330.62
	12	74	77	2520.16	76	-74374.78
	13	74	76	2522.08	76	-74374.78
N.I.	1.0	100	1000	- A	1	7C I .

No model exists between  $\underline{p}=74$  and  $\overline{p}=76$ , so algo stops and returns the simpler model with 74 peaks.

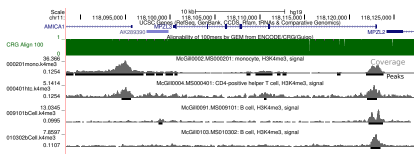
### Chromatin immunoprecipitation sequencing (ChIP-seq)

Analysis of DNA-protein interactions.



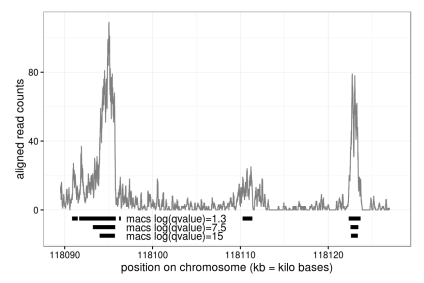
Source: "ChIP-sequencing," Wikipedia.

#### Problem: find peaks in each of several samples

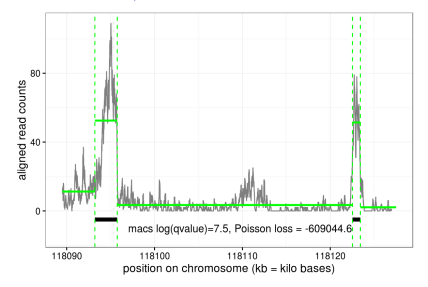


- Grey profiles are noisy aligned read count signals peaks are genomic locations with protein binding sites.
- ▶ Black bars are peaks called by MACS2 (Zhang et al, 2008) many false positives! (black bars where there is only noise)
- ► From a machine learning perspective, this is binary classification (positive=peaks, negative=noise).

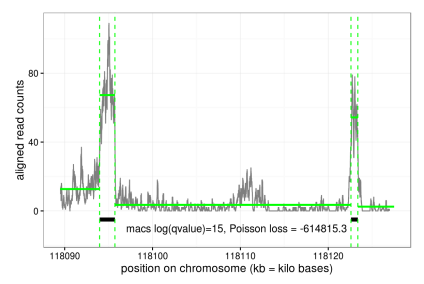
#### Which macs parameter is best for these data?



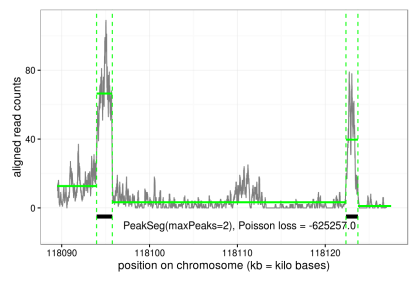
#### Compute likelihood/loss of piecewise constant model



#### Idea: choose the parameter with a lower loss



#### PeakSeg: search for the peaks with lowest loss



Simple model with only one parameter (number of peaks).



#### Statistical model is a piecewise constant Poisson mean

Het We have in 200 up t data  $z_1, \ldots, z_n \in \mathbb{Z}_+$ .

- ▶ Fix the number of segments  $S \in \{1, 2, ..., n\}$ .
- ▶ Optimization variables: S-1 changepoints  $t_1 < \cdots < t_{S-1}$  and S segment means  $u_1, \ldots, u_S \in \mathbb{R}_+$ .
- ▶ Let  $0 = t_0 < t_1 < \cdots < t_{S-1} < t_S = n$  be the segment limits.
- Statistical model: for every segment  $s \in \{1, ..., S\}$ ,  $z_i \stackrel{\text{iid}}{\sim} \mathsf{Poisson}(u_s)$  for every data point  $i \in (t_{s-1}, t_s]$ .
- PeakSeg up-down constraint:  $u_1 \le u_2 \ge u_3 \le u_4 \ge \cdots$
- Want to find means  $u_s$  which maximize the Poisson likelihood:  $P(Z = z_i | u_s) = u_s^{z_i} e^{-u_s} / (z_i!)$ .
- Equivalent to finding means  $u_s$  which minimize the Poisson loss:  $\ell(u_s, z_i) = u_s z_i \log u_s$ .

#### Dynamic programming and functional pruning

Classical dynamic programming for optimal partitioning (Jackson et al 2005) computes the vector of optimal loss values up to N data points,  $O(N^2)$  time because each DP iteration needs to consider all O(N) possible changepoints and cost values.

$$C_1$$
  $C_2$   $\cdots$   $C_{N-1}$   $C_N$ 

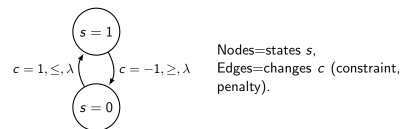
Functional pruning optimal partitioning (Maidstone 2016) computes a vector of loss functions,  $O(N \log N)$  because each DP iteration only considers  $O(\log N)$  candidate changepoints (the others — which will never be optimal — are pruned).

$$C_1(m_1)$$
  $C_2(m_2)$  ···  $C_{N-1}(m_{N-1})$   $C_N(m_N)$ 

**Contribution of this work**: a new algorithm that applies the functional pruning technique to the up-down constrained model.

#### Constrained optimal partitioning problem

$$\begin{aligned} & \underset{\mathbf{c} \in \mathbb{R}^N, \ \mathbf{s} \in \{0,1\}^N}{\text{minimize}} & & \sum_{i=1}^N \ell(m_i, z_i) + \lambda \sum_{i=1}^{N-1} I(c_i \neq 0) \\ & \text{subject to} & \text{no change: } c_t = 0 \Rightarrow m_t = m_{t+1} \text{ and } s_t = s_{t+1} \\ & \text{go up: } c_t = 1 \Rightarrow m_t \leq m_{t+1} \text{ and } (s_t, s_{t+1}) = (0, 1), \\ & \text{go down: } c_t = -1 \Rightarrow m_t \geq m_{t+1} \text{ and } (s_t, s_{t+1}) = (1, 0). \end{aligned}$$



# Generalized Functional Pruning Optimal Partitioning (GFPOP) algorithm for up-down constrained model

Recursively compute two vectors of real-valued cost functions:

$$\overline{C}_1(m_1)$$
  $\cdots$   $\overline{C}_N(m_N)$  optimal cost in peak state  $s=1$   $\underline{C}_1(m_1)$   $\cdots$   $\underline{C}_N(m_N)$  optimal cost in background state  $s=0$ 

$$\overline{C}_{t+1}(\mu) = \ell(\mu, z_i) + \min\{\overline{C}_t(\mu), \underline{C}_t^{\leq}(\mu) + \lambda\},$$

$$\underline{C}_{t+1}(\mu) = \ell(\mu, z_i) + \min\{\underline{C}_t(\mu), \overline{C}_t^{\geq}(\mu) + \lambda\},$$

$$\text{where } f^{\leq}(\mu) = \min_{x \leq \mu} f(x),$$

$$f^{\geq}(\mu) = \min_{x \geq \mu} f(x).$$

$$\underline{C}_t$$

$$\underline{C}_t$$

$$\underline{C}_t$$