다변량 자료분석

Choosing Starting Values

for the EM algorithm

in Multivariate Gaussian Mixture Models

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## 논문 소개

제목 Choosing starting values for the EM algorithm for getting the highest likelihood in

multivariate Gaussian mixture models

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### Mixture Model & EM algorithm

- EM 알고리즘은 초기값에 매우 민감하다.
- 특히 다변량 자료를 이용하는 경우 초기값의 영향이 매우 크다.
- Mixture model 추정에서 likelihood의 local maximum을 선택할 위험이 있다.
   ex) cluster의 개수를 잘못 추정할 수 있음
- → 제한된 반복횟수 안에서 적절한 초기값을 선택하는 몇 가지 방법을 제안 & 비교

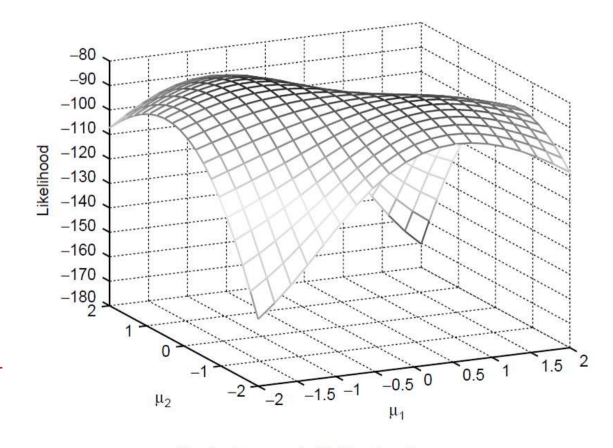


Fig. 1. A two-mode likelihood surface.

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#### Gaussian Mixture Model

 $\mathbf{x}_1, \dots, \mathbf{x}_n$  in  $\mathbf{R}^d$  from a random vector with density

$$f(\mathbf{x}) = \sum_{k=1}^{K} p_k \phi(\mathbf{x} | \boldsymbol{\mu}_k, \boldsymbol{\Sigma}_k)$$

 $p_k$ : mixing proportions, k = 1, ..., K

 $\phi(\mathbf{x}|\boldsymbol{\mu},\boldsymbol{\Sigma})$ : multivariate Normal density with mean vector  $\boldsymbol{\mu}$  and variance matrix  $\boldsymbol{\Sigma}$ 

### ML Estimateion using the EM algorithm

 $\theta = (p_1, ..., p_K, \mu_1, ..., \mu_K, \Sigma_1, ... \Sigma_K)$  are estimated by maximizing the log-likelihood

$$l(\theta|\mathbf{x}_1, ..., \mathbf{x}_n) = \sum_{i=1}^n \ln \left[ \sum_{k=1}^K p_k \phi(\mathbf{x}_i|\boldsymbol{\mu}_k, \boldsymbol{\Sigma}_k) \right]$$

Starting from an initial parameter  $\theta^0$ , iterate the following E-M steps:

**E step** Compute  $\hat{p}_k(\mathbf{x}_i)$  which are the current conditional probabilities that  $\mathbf{x}_i$  from the kth cluster

**M step** Update the ML estimates  $(\hat{p}_k, \hat{\mu}_k, \hat{\Sigma}_k)$  using  $\hat{p}_k(\mathbf{x}_i)$  as conditional mixing weights

# Methods for Choosing Starting Values

- 1. Short runs of EM
- 2. CEM (Classification EM)
- 3. SEM (Stochastic EM)

### CEM algorithm (Classification EM)

**E step** Compute  $\hat{p}_k(\mathbf{x}_i)$  which are the current conditional probabilities that  $\mathbf{x}_i$  from the kth cluster

**C step** Design a partition  $P = (P_1, ..., P_K)$  of  $\mathbf{x}_1, ..., \mathbf{x}_n$  by assigning  $\mathbf{x}_i$  to the cluster maximizing  $\hat{p}_k(\mathbf{x}_i)$ 

**M step** Compute the ML estimates  $(\hat{p}_k, \hat{\mu}_k, \hat{\Sigma}_k)$  using the cluster  $P_k$  as sub-sample of the kth cluster

It is a *K-means*-like algorithm, converges in a finite number of iterations.

When  $z_i$  is the missing cluster label of  $\mathbf{x}_i$ , it maximize the complete data log-likelihood

$$Cl(\theta|z_1,\ldots,z_n,\mathbf{x}_1,\ldots,\mathbf{x}_n) = \sum_{k=1}^K \sum_{\{i:z_i=k\}} \ln[p_k \phi(\mathbf{x}_i|\boldsymbol{\mu}_k,\boldsymbol{\Sigma}_k)]$$

### SEM algorithm (Stochastic EM)

**E step** Compute  $\hat{p}_k(\mathbf{x}_i)$  which are the current conditional probabilities that  $\mathbf{x}_i$  from the kth cluster

**S step** Design a partition  $P = (P_1, ..., P_K)$  of  $\mathbf{x}_1, ..., \mathbf{x}_n$  by assigning  $\mathbf{x}_i$  at random to one of the clusters according to the Multinomial distribution with parameter  $\hat{p}_k(\mathbf{x}_i)$ 

**M step** Compute the ML estimates  $(\hat{p}_k, \hat{\mu}_k, \hat{\Sigma}_k)$  using the cluster  $P_k$  as sub-sample of the kth cluster

### SEM algorithm (Stochastic EM)

It is same as the Monte Carlo EM algorithm with a single replication, and generates Markov chain.

Parameter estimate from a SEM sequence  $(\theta^r)_{r=1,\dots,R}$ :

1. The mean value of the sequence after a burn-in period b

$$\hat{\theta} = \sum_{r=b+1}^{R} \theta^r / (R - b)$$

2. The value leading to the highest likelihood in the sequence

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# Methods for Choosing Starting Values

- 1. Random initialization
- 2. Using short runs of EM
- 3. Using the CEM algorithm
- 4. Using the SEM algorithm

# Methods for Choosing Starting Values

1. Random initialization	EM
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- 2. Using short runs of EM **em-EM**
- 3. Using the CEM algorithm **CEM-EM**
- 4. Using the SEM algorithm **SEM-EM**

#### **Experimental Strategies**

- 1. EM, CEM, SEM 알고리즘을 1번 반복할 때 계산 시간에 큰 차이가 없다.
  - → 따라서 총 반복수를 동일하게 설정하여 총 계산 시간을 제한
- 2. 동일한 총 반복수 하에 각 알고리즘을 10번 반복하는 방법을 포함하여 총 8개의 방법을 비교 (iteration) (repetition)

#### EX. Total Number of Iterations = 1000

<b>1EM</b> 1000 iteration	ons for EM
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**10EM** 10 repetitions of 100 iterations for each EM run

**1em-EM** 500 iterations for em and 500 iterations for EM

**10em-EM** 10 repetitions of 50 iterations for em and 50 iterations for EM

**1CEM-EM** 500 iterations for CEM and 500 iterations for EM

**10CEM-EM** 10 repetitions of 50 iterations for CEM and 50 iterations for EM

**SEMmean-EM** & **SEMmax-EM** 500 iterations for SEM and 500 iterations for EM

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Data set	P1	P2	P3	P1 noise, P2 noise, P3 noise
Dimension	d=2	d=2	d = 2	
Number of Clusters	K = 2	K = 2	K = 4	
Mixing Proportion	$p_1 = p_2 = 0.5$	$p_1 = 0.7, p_2 = 0.3$	$p_1 = p_2 = p_3 = p_4 = 0.5$	Add noise to P1, P2, P3
Parameters	$\mu_1 = (0,0)'$ $\mu_2 = (2.5,0)'$ $\operatorname{diag}(\mathbf{\Sigma}_1) = \left(3, \frac{1}{3}\right)$ $\operatorname{diag}(\mathbf{\Sigma}_2) = \left(\frac{1}{3}, 3\right)$	$\mu_1 = \mu_2 = (0,0)'$ $\operatorname{diag}(\mathbf{\Sigma}_1) = \left(3, \frac{1}{3}\right)$ $\operatorname{diag}(\mathbf{\Sigma}_2) = \left(\frac{1}{3}, 3\right)$	$\mu_{1} = (0, -2)', \ \mu_{2} = (2.0)'$ $\mu_{3} = (0, 2)', \ \mu_{4} = (-2, 0)'$ $\operatorname{diag}(\mathbf{\Sigma}_{1}) = \operatorname{diag}(\mathbf{\Sigma}_{2}) = \left(3, \frac{1}{3}\right)$ $\operatorname{diag}(\mathbf{\Sigma}_{3}) = \operatorname{diag}(\mathbf{\Sigma}_{4}) = \left(\frac{1}{3}, 3\right)$	from $Uniform$ $[-0.8, 0.8] \times [-0.8, 0.8]$ with proportion 0.2

총 반복수 = (60, 120, 240, 480, …, 15360)

※ 총 반복수 960을 기준으로 Small / Large로 구분

**Table1** Small v.s. Large

	EM		CEM-	EM	em-EN	1	SEM-EM	
nb. it.	1	10	1	10	1	10	Mean	Max
Small	7	0	6	4	0	0	6	4
Large	35	98	33	87	45	98	46	41

Table2
Single run
v.s.
Repeated runs

	EM		CEM-	ЕМ	em-EN	1	SEM-EM	
nb. it.	1	10	1	10	1	10	Mean	Max
Small	79	21	67	29	88	12	11	46
Large	5	36	5	20	10	12	0	6

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iterations

**Table3**Comparison by pairs with a large number of

	P1	P1 noise	P2	P2 noise	P3	P3 noise
10EM vs. 10CEM-EM	2-0	38-27	19-0	30-42	36-0	3-86
10EM vs. 10em-EM	0-0	8-28	1-0	7-43	5-1	39-39
10EM vs. SEMmax-EM	0-0	43-8	0-0	43-21	5-3	23-64
10CEM-EM vs. 10em-EM	0-2	6-40	0-19	2-30	1-35	83-4
10CEM-EM vs. SEMmax-EM	0-2	54-30	0-19	61-21	0-35	66-8
10em-EM vs. SEMmax-EM	0-0	57-10	0-1	63-6	5-7	29-56

Table4
Means & standard
deviations of maximum
log-likelihood

	P1	P1 noise	P2	P2 noise	P3	P3 noise
10EM	-659.8	-909.2	-616.1	-881.7	-754.3)	-928.2
	(14.6)	(13.1)	(17.8)	(17.3)	(13.2)	(13.7)
10CEM-EM	-659.8	-909.9	-617.9	-881.2	-755.6	-919.8
	(14.6)	(12.5)	(18.9)	(18.4)	(13.3)	(12.3)
10em-EM	-659.8	-908.3	-616.1	-880.2	-754.3	-927.4
	(14.6)	(12.3)	(17.8)	(17.6)	(13.2)	(14.0)
SEMmax-EM	-659.8	-911.1	-616.1	-883.7	-754.3	-925.5
	(14.6)	(13.9)	(17.8)	(17.8)	(13.2)	(13.0)



Noise가 없는 data에서는 10CEM-EM이 가장 안 좋지만 cluster의 개수가 적을 때는 차이가 크지 않다. Noisy data에서는 결론이 명확하지 않다.

## Real Data Sets

Data set	Stars	Geyser	Biological Profiles
Number of Observations	n = 2370	n = 272	n = 3641
Dimension	d = 2	d = 2	d = 5
Number of Clusters	K = 3	K = 3	K = 10
Explanation	Stars described by their velocity <i>U</i> toward the galactic center and velocity <i>V</i> toward the galactic rotation (Soubiran, 1993)	Eruptions of the Old Faithful Geyser in Yellowstone National Park measured by the <i>duration</i> of the eruption, and the <i>waiting time</i> before the next eruption (Venables and Ripley, 1994)	Biological profiles of patients (Sandor, 1976)

### Real Data Sets - Stars & Geyser

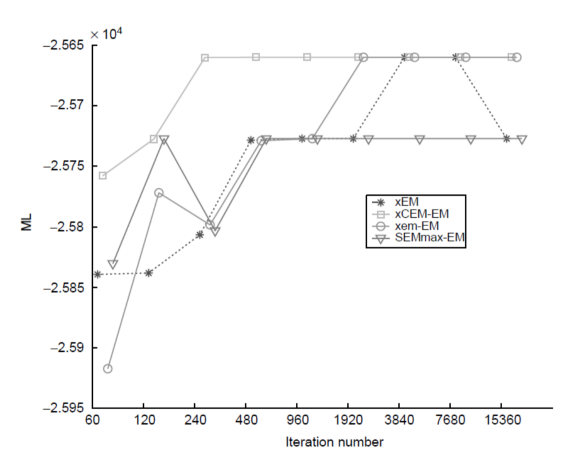


Figure 1. Stars (x=10)

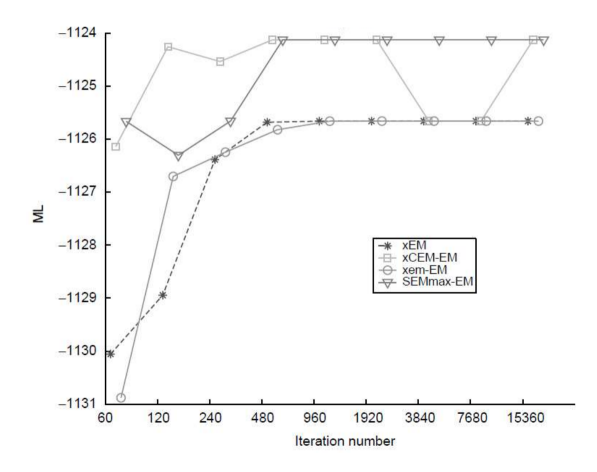


Figure 2. Geyser (x=10)

### Real Data Sets - Stars & Geyser

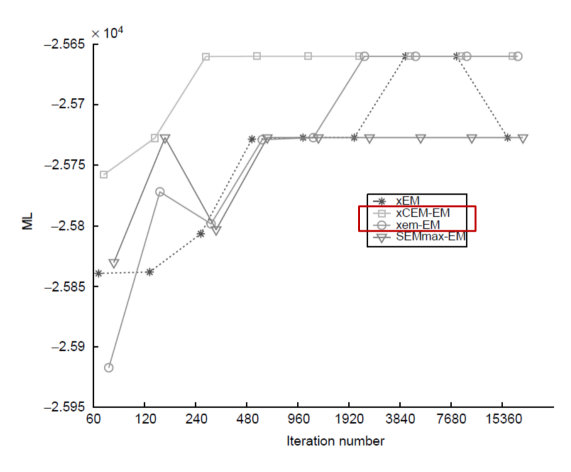


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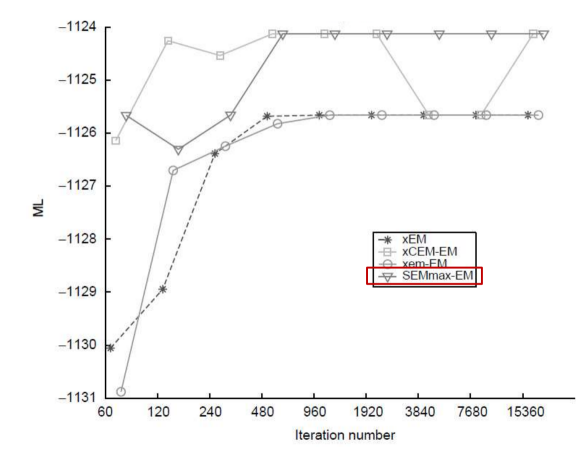
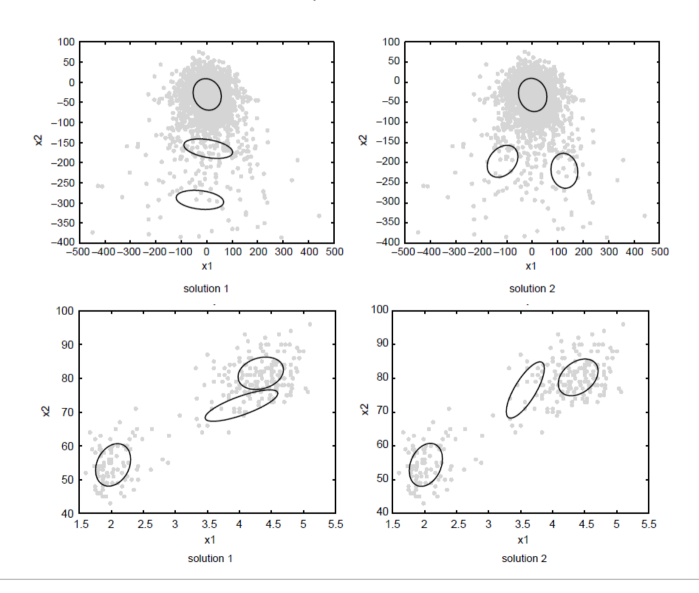


Figure 2. Geyser (x=10)

#### Real Data Sets - Stars & Geyser



- < Two solutions in competition >
  They provide isodensity ellipses for each cluster.
- Solution 1 provides the highest likelihood.

Figure 3. Solutions of data set "Stars"

Figure 4. Solutions of data set "Geyser"

# Real Data Sets - Biological Profiles

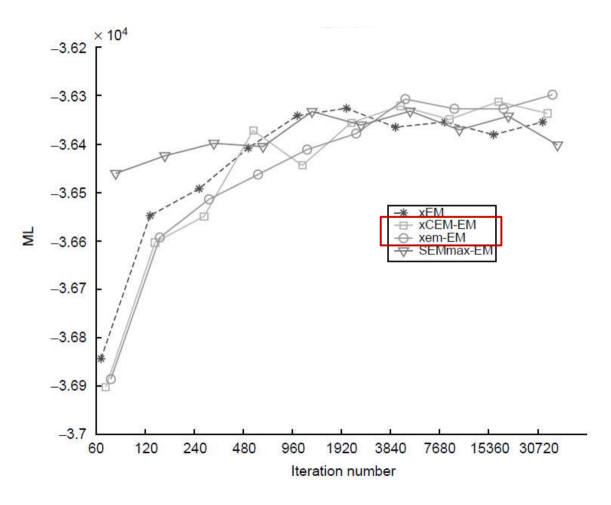


Figure 5. Biological Profiles (x=10)

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#### Discussion

- 1. 총 반복수가 충분해야 한다.
- 2. 알고리즘을 여러 번 반복 (repetition) 하는 방법이 좋다. (ex. 10em-EM)
- 3. short runs of EM 방법의 성능이 다른 방법들에 비해 약간 더 높으며, 기본적인 EM 방법이 가장 나쁘다.
- 4. short runs of EM 방법은 단순하고 특별한 가정이 필요하지 않으며, noisy data에 덜 민감하기 때문에 CEM 또는 SEM 알고리즘보다 더 적절하다.
- 5. 결론적으로 총 반복수가 충분히 크다면, short runs of EM의 반복을 통해 초기값을 선택하는 것을 추천한다.

# 사이 감사합니다