# Spectral Unmixing: Estimating partial abundances

Pravesh Debba

CSIR Built Environment Logistics and Quantitative Methods (LQM)

LQM Seminar Presentation 2009

Debba (CSIR)

Spectral Unmixing

LQM 2009 1 / 22

(3)

# Outline

## 1 Background and Research Question

- 2 Method of spectral unmixing
- 3) End-member spectra and synthetic mixtures

## 4 Results



イロト イ団ト イヨト イヨト

# Outline





End-member spectra and synthetic mixtures

#### 4 Results



Debba (CSIR)



- Background and Research Question
- 2 Method of spectral unmixing
- 3 End-member spectra and synthetic mixtures







- Background and Research Question
- 2 Method of spectral unmixing
- 3 End-member spectra and synthetic mixtures





- Background and Research Question
- 2 Method of spectral unmixing
- 3 End-member spectra and synthetic mixtures





# If research could be as easy as eating a chocolate cake ....



Figure: Can you guess the ingredients for this chocolate cake?

Ingredients	Quantity
unsweetened chocolate	
unsweetened cocoa powder	
boiling water	
flour	
baking powder	
baking soda	
salt	
unsalted butter	
white sugar	
eggs	
pure vanilla extract	
milk	

## Table: Chocolate cake ingredients

イロト イヨト イヨト イヨト

Ingredients	Quantity
unsweetened chocolate	120 grams
unsweetened cocoa powder	28 grams
boiling water	240 ml
flour	315 grams
baking powder	2 teaspoons
baking soda	1 teaspoon
salt	1/4 teaspoon
unsalted butter	226 grams
white sugar	400 grams
eggs	3 large
pure vanilla extract	2 teaspoons
milk	240 ml

Table: Chocolate cake ingredients

(日) (四) (日) (日) (日)

# What is spectral unmixing?



Figure: The concept of unmixing - taken from
http://www.emeraldinsight.com/fig/0830140104006.png

LQM 2009 6 / 22

- Most spectral unmixing techniques are variants of algorithms involving matrix inversion.
- Major problem in spectral unmixing is the non-orthogonality of end-members.
- Ability to estimate abundances in complex mixtures through spectral unmixing techniques – complicated when considering very similar spectral signatures.
- Iron-bearing oxide/hydroxide/sulfate minerals have similar spectral signatures.

How could estimates of abundances of spectrally similar iron-bearing oxide/hydroxide/sulfate minerals in complex mixtures be obtained using hyperspectral data?

- Most spectral unmixing techniques are variants of algorithms involving matrix inversion.
- Major problem in spectral unmixing is the non-orthogonality of end-members.
- Ability to estimate abundances in complex mixtures through spectral unmixing techniques – complicated when considering very similar spectral signatures.
- Iron-bearing oxide/hydroxide/sulfate minerals have similar spectral signatures.

How could estimates of abundances of spectrally similar iron-bearing oxide/hydroxide/sulfate minerals in complex mixtures be obtained using hyperspectral data?

- Most spectral unmixing techniques are variants of algorithms involving matrix inversion.
- Major problem in spectral unmixing is the non-orthogonality of end-members.
- Ability to estimate abundances in complex mixtures through spectral unmixing techniques – complicated when considering very similar spectral signatures.
- Iron-bearing oxide/hydroxide/sulfate minerals have similar spectral signatures.

How could estimates of abundances of spectrally similar iron-bearing oxide/hydroxide/sulfate minerals in complex mixtures be obtained using hyperspectral data?

- Most spectral unmixing techniques are variants of algorithms involving matrix inversion.
- Major problem in spectral unmixing is the non-orthogonality of end-members.
- Ability to estimate abundances in complex mixtures through spectral unmixing techniques – complicated when considering very similar spectral signatures.
- Iron-bearing oxide/hydroxide/sulfate minerals have similar spectral signatures.

How could estimates of abundances of spectrally similar iron-bearing oxide/hydroxide/sulfate minerals in complex mixtures be obtained using hyperspectral data?

- Most spectral unmixing techniques are variants of algorithms involving matrix inversion.
- Major problem in spectral unmixing is the non-orthogonality of end-members.
- Ability to estimate abundances in complex mixtures through spectral unmixing techniques – complicated when considering very similar spectral signatures.
- Iron-bearing oxide/hydroxide/sulfate minerals have similar spectral signatures.

How could estimates of abundances of spectrally similar iron-bearing oxide/hydroxide/sulfate minerals in complex mixtures be obtained using hyperspectral data?

۲

- **(() ) ) ( () ) ) () )** 

# Old method: problem

Linear Spectral Mixture Analysis (LSMA): The observed spectrum U for any given pixel in the scene is expressed as:

$$U = Rp + \epsilon$$
 where  $\sum_{i=1}^{n} p_i = 1$  and  $0 \le p_i \le 1$ 

and R is a matrix of which each column corresponds to an endmember, p is a column vector that denotes the abundances and  $\epsilon$  denotes the residual term.

Minimize:  $\sum_{i=1}^{n} \epsilon_i^2 = \sum_{i=1}^{n} \left( \sum_{j=1}^{m} (R_{i,j} \times p_i) - U_i \right)^2$ 

Solution:  $\hat{p} = \left(R^T R\right)^{-1} R^T U$ 

# New method: solution

Suppose *M* is an exhaustive set of endmembers and  $E \subset M$  is a set of endmembers under consideration for unmixing. Each component spectrum  $e \in E$  consists of *L* discrete wavelengths  $\lambda_I$  (I = 1, ..., L). It is denoted by  $R^e = (R^e(\lambda_1), ..., R^e(\lambda_L))$ , where  $R^e(\lambda_I)$  is the reflectance value at wavelength  $\lambda_I$ .

The observed spectrum U for any given pixel in the scene is expressed as:

$$U = \begin{bmatrix} R^{E} \\ R^{M \setminus E} \end{bmatrix} \times (p^{E} p^{M \setminus E}) + \epsilon \text{ where } \sum_{e=1}^{||E||} p_{e} \leq 1 \text{ and } 0 \leq p_{e} \leq 1$$

Accordingly, a spectrum at  $\lambda_I$  can be modeled as

$$\widehat{U}(\lambda_l) = \sum_{e=1}^{||\mathcal{E}||} p_e R^e(\lambda_l) + p_0 R^{M \setminus \mathcal{E}}(\lambda_l), \qquad (1)$$

where  $0 \le p_e \le 1$ ,  $p_0 + \sum_{e=1}^{||E||} p_e = 1$  and  $0 \le p_0 \le 1$ , so the set of  $p_0 \ge 1$ .

The difference between the estimated and actual spectra at  $\lambda_I$  equals

$$\omega_l = U(\lambda_l) - \sum_{e=1}^{||E||} p_e R^e(\lambda_l) \,. \tag{2}$$

Minimization of some function of  $\omega_l$ , e.g. SumSpec =  $\sum_{l=1}^{L} |\omega_l|$  or VarSpec = var( $\omega_l$ ) results in estimates for  $p_e$ .

Alternatively: Use either the differences in the first derivative or the second derivative instead of the actual differences. The difference in the first derivative between an estimated and an actual spectrum at  $\lambda_I$  is

$$\omega_{I}^{\prime} = \frac{\Delta U(\lambda_{I})}{\Delta \lambda_{I}} - \sum_{e=1}^{||E||} p_{e} \left(\frac{\Delta R^{e}(\lambda_{I})}{\Delta \lambda_{I}}\right), \qquad (3)$$

where  $\Delta x_l = x_{l+1} - x_l$ . Minimization of a loss function of equation 3, e.g. SumDeriv =  $\sum_{l=1}^{L-2} |\omega'_l|$  or VarDeriv =  $var(\omega'_l)$ , results in estimates of  $p_e$ . The minimization is achieved through simulated annealing, using either SumSpec, SumDeriv, VarSpec or VarDeriv as the fitness function to optimize.

Debba (CSIR)

Simulated annealing is a general optimization method of a fitness function  $\phi(\omega)$  – depends on  $p_e$ . Starting with a random configuration of  $p_e$ ,  $\phi(\omega^0)$ is calculated. Let  $\omega^i$  and  $\omega^{i+1}$  represent two solutions with fitness  $\phi(\omega^i)$ and  $\phi(\omega^{i+1})$ . Configuration  $\omega^{i+1}$  is derived from  $\omega^i$  by randomly replacing one point  $p_j$  of  $\omega^i$  by a new point  $p_k$  in  $\left|0, 1 + p_j - \sum p_e\right|$ , so that  $\sum p_e < 1$ . A probabilistic acceptance criterion decides whether  $\omega^{i+1}$  is accepted or not i.e.

$$P_{c}(\omega^{i} \to \omega^{i+1}) = \begin{cases} 1, & \text{if } \phi(\omega^{i+1}) \leq \phi(\omega^{i}) \\ \exp\left(\frac{\phi(\omega^{i}) - \phi(\omega^{i+1})}{\mathbf{c}}\right), & \text{if } \phi(\omega^{i+1}) > \phi(\omega^{i}) \end{cases}$$
(4)

where  $\mathbf{c}$  denotes a parameter and is reduced by a factor of 0.95, thereby decreasing the probability of accepting inferior moves. Reduction stops when the process stabilizes. A transition takes place if  $\omega^{i+1}$  is accepted. Next, a solution  $\omega^{i+2}$  is derived from  $\omega^{i+1}$ , and the probability  $P_{c}(\omega^{i+1} \rightarrow \omega^{i+2})$  is calculated with a similar acceptance criterion as equation 4. The fitness function will be one of SumSpec, VarSpec, SumDeriv or VarDeriv. イロト 不得 トイヨト イヨト 二日

Debba (CSIR)

LQM 2009 11 / 22



Figure: Five end-members spectra from USGS library, resampled to DAIS VIR region

Debba (CSIR)

LQM 2009 12 / 22



Figure: Five end-members spectra with error from the U(-0.02, 0.02) distribution. Smoothing was applied to the spectra.



Figure: Mixed spectra with error from the U(-0.02, 0.02) distribution. Smoothing was applied to the spectra.

LQM 2009 14 / 22



Figure: First derivative of end-member spectra after applying smoothing.

Debba (CSIR)

LQM 2009 15 / 22

End-member spectra and synthetic mixtures



LQM 2009 16 / 22

Results										
	Table: Abundance: Using observed spectra & SumSpec.									
Known abundance						Estima	ted abi	undanc	e	
	Goe	Jar	Сор	Fer	Goe	Jar	Сор	Fer	$M \setminus E$	
	End-member spectrum included in E									
	1.00	0.00	0.00	0.00	0.78	0.03	0.05	0.14	0.00	
	0.00	1.00	0.00	0.00	0.03	0.94	0.00	0.01	0.02	
	0.00	0.00	1.00	0.00	0.03	0.06	0.89	0.02	0.00	
	0.00	0.00	0.00	1.00	0.03	0.00	0.01	0.92	0.04	
	End-r	nember	spectr	um exc	cluded f	rom E				
	1.00	0.00	0.00	0.00	—	0.25	0.00	0.64	0.11	
	0.00	1.00	0.00	0.00	0.00	—	1.00	0.00	0.00	
	0.00	0.00	1.00	0.00	0.01	0.68	—	0.00	0.31	
	0.00	0.00	0.00	1.00	0.71	0.00	0.00	—	0.29	
	Mixtu	ires		I						
	0.50	0.50	0.00	0.00	0.48	0.51			0.01	
	0.50	0.50	0.00	0.00	0.40	0.52	0.01	0.05	0.02	
	0.15	0.25	0.25	0.35	0.66	0.30			0.04	
	0.15	0.25	0.25	0.35	0.20	0.26	0.23	0.29	0.02	

Debba (CSIR)

LQM 2009 17 / 22

Results										
	Table: Abundance: Using observed spectra & VarSpec.									
Known abundance						Estima	ted abi	undanc	e	
	Goe	Jar	Сор	Fer	Goe	Jar	Сор	Fer	$M \setminus E$	
	End-member spectrum included in <i>E</i>									
	1.00	0.00	0.00	0.00	0.90	0.02	0.00	0.08	0.00	
	0.00	1.00	0.00	0.00	0.07	0.91	0.00	0.00	0.02	
	0.00	0.00	1.00	0.00	0.02	0.01	0.94	0.00	0.03	
	0.00	0.00	0.00	1.00	0.03	0.00	0.01	0.92	0.04	
	End-r	nember	spectr	rum exc	cluded f	rom E				
	1.00	0.00	0.00	0.00	—	0.26	0.00	0.69	0.05	
	0.00	1.00	0.00	0.00	0.43		0.57	0.00	0.00	
	0.00	0.00	1.00	0.00	0.01	0.38	—	0.00	0.61	
	0.00	0.00	0.00	1.00	0.97	0.00	0.01	—	0.02	
	Mixtu	ires								
	0.50	0.50	0.00	0.00	0.49	0.50			0.01	
	0.50	0.50	0.00	0.00	0.35	0.54	0.01	0.09	0.01	
	0.15	0.25	0.25	0.35	0.55	0.24	—		0.21	
	0.15	0.25	0.25	0.35	0.14	0.25	0.23	0.34	0.04	

Debba (CSIR)

LQM 2009 18 / 22

Results										
Table: Abundance: Using 1st derivative spectra & SumDeriv.										
Kr	nown a	bundan	ce		Estima	ated ab	undanc	e		
Goe	Jar	Сор	Fer	Goe	Jar	Сор	Fer	$M \setminus E^1$		
End-r	End-member spectrum included in <i>E</i>									
1.00	0.00	0.00	0.00	0.87	0.02	0.00	0.06	0.05		
0.00	1.00	0.00	0.00	0.00	0.94	0.06	0.00	0.00		
0.00	0.00	1.00	0.00	0.01	0.04	0.91	0.00	0.05		
0.00	0.00	0.00	1.00	0.07	0.01	0.00	0.86	0.06		
End-r	nembei	specti	rum exc	cluded f	rom E					
1.00	0.00	0.00	0.00		0.33	0.00	0.66	0.01		
0.00	1.00	0.00	0.00	0.48	—	0.51	0.00	0.01		
0.00	0.00	1.00	0.00	0.00	0.45	—	0.01	0.54		
0.00	0.00	0.00	1.00	0.33	0.00	0.00		0.67		
Mixtu	ires									
0.50	0.50	0.00	0.00	0.48	0.51			0.01		
0.50	0.50	0.00	0.00	0.46	0.48	0.05	0.01	0.00		
0.15	0.25	0.25	0.35	0.27	0.32	—		0.41		
0.15	0.25	0.25	0.35	0.09	0.24	0.26	0.39	0.02		

LQM 2009 19 / 22

Results										
Table: Abundance: Using 1st derivative spectra & VarDeriv.										
Kr	nown a	bundan	ce		Estima	ated ab	undand	ce		
Goe	Jar	Сор	Fer	Goe	Jar	Сор	Fer	$M \setminus E^1$		
End-r	End-member spectrum included in <i>E</i>									
1.00	0.00	0.00	0.00	0.90	0.00	0.03	0.07	0.00		
0.00	1.00	0.00	0.00	0.01	0.92	0.04	0.01	0.02		
0.00	0.00	1.00	0.00	0.01	0.07	0.92	0.00	0.00		
0.00	0.00	0.00	1.00	0.08	0.00	0.00	0.89	0.03		
End-r	nembei	specti	rum exc	cluded f	from E					
1.00	0.00	0.00	0.00		0.28	0.00	0.56	0.16		
0.00	1.00	0.00	0.00	0.31		0.68	0.00	0.01		
0.00	0.00	1.00	0.00	0.00	0.46	—	0.00	0.54		
0.00	0.00	0.00	1.00	0.27	0.00	0.00	—	0.73		
Mixtu	ires									
0.50	0.50	0.00	0.00	0.44	0.53			0.03		
0.50	0.50	0.00	0.00	0.45	0.52	0.02	0.01	0.00		
0.15	0.25	0.25	0.35	0.05	0.40			0.55		
0.15	0.25	0.25	0.35	0.07	0.27	0.24	0.37	0.05		

LQM 2009 20 / 22

Table: Correlation coefficient between pairs of original spectra, pairs of first derivative of spectra and pairs of second derivative of spectra.

Original spectra:									
	goethite	jarosite	copiapite	ferrihydrite					
goethite	1.00								
jarosite	0.67	1.00							
copiapite	0.43	0.72	1.00						
ferrihydrite	0.86	0.29	0.16	1.00					
1st derivativ	re (lower ∆	) & 2nd d	lerivative (u	pper $ riangle$ ):					
	goethite	jarosite	copiapite	ferrihydrite					
goethite	1.00	0.35	-0.14	0.22					
jarosite	0.71	1.00	0.43	0.18					
copiapite	0.35	0.79	1.00	-0.02					
ferrihydrite	0.44	0.24	-0.15	1.00					

LQM 2009 21 / 22

< ロ > < 同 > < 三 > < 三

This study resulted in four main conclusions.

- Abundances of spectrally similar minerals in mine wastes can be estimated with relatively high accuracy by unmixing of first derivatives of target spectra, in which contributing components are decorrelated.
- Simulated annealing proved efficient in minimizing variance of the difference spectrum to estimate abundance of spectrally similar minerals.
- Higher accuracy of abundance estimates is gained when end-member spectra contributing to target spectra is included.
- The choice to use the original spectra, the first or second derivatives spectra depends on the signal-to-noise ratio of the sensor device. Higher signal-to-noise ratios allows better accuracy in the abundance estimation by using higher order derivatives.

イロト イポト イヨト イヨト 二日