

Are Silanol Groups Bad or Good for Basic Compounds?

Sunnise C18 Sunnise C18-SAC Silanol Activity Controlled C18 Column

# New-Type RP Column



ChromaNik Technologies Inc. Email: info@chromanik.co.jp http://chromanik.co.jp

Sunrise C18 and C18-SAC Silanol Activity Controlled C18 HPLC Column



## New generation reversed-phase utilized silanol groups

### Silanol group and peak tailing

It is generally said that residual silanol groups on a stationary phase such as C18 (ODS) causes absorption or peak tailing for a sample. Especially silanol groups near a hydrophobic site don't solvate with water completely, so that they show high absorption for basic compounds. Its peak shows terribly tailing. Several endcapping techniques have been developed to solve these problems for many years.

### Silanol activity control technology

ChromaNik developed the technique that decreased only silanol groups with high absorption activity to a basic compound and remained effective sailnol groups on the stationary phase. Silanol activity control and no end-capping led the existence of silanol groups with high hydration which created a new and unique reversed-phase separation mode including hydrogen bond and ion-exchange interaction. Furthermore, silanol activity controlling, then end-capping technique improved a peak shape of a basic compound exceedingly.





# Feature of Sunrise series

### Sunrise C18

- •The "1st Choice" column as a fully end-capped C18 column
- •Full end-capping after silanol activity control
- Reducing adsorption of a basic compound extremely
- A good peak shape for a metal cheleting compound
- Widely available for general reversed-phase separation

### Sunrise C18-SAC

- •The "2nd Choice" column which takes advantage of effective silanol groups interaction
- Reducing silanol groups with high adsorption activity
- •The new separation mechanism including hydrogen bond and ion-exchange interaction
- Effective for separation of a basic compound and a polar compound
- Different selectivity and improvement of separation without changing a mobile phase

#### The elution order of pyridine



Sunrise C18 and C18-SAC



Silanol Activity Controlled C18 HPLC Column

# Sunrise series create an unique separation

# \* Effectiveness of silanol activity control: Comparison between Sunrise C18 and C18-SAC

Sunrise C18 is the so-called fully end-capped C18 column. It shows the same separation behavior as a conventional C18 column.

On the other hand, Sunrise C18-SAC shows hydrogen bond and ion-exchange interactions based on a residual silanol on the silica support in addition to reversed-phase separation. For example Sunrise C18 column separates a basic compound similarly as a conventional C18, while Sunrise C18-SAC makes retention of a basic compound be large because an ion-exchange interaction works although a non-ionic compound shows the almost same retention on both Sunrise C18 and C18-SAC. Furthermore, Sunrise C18-SAC shows large retention for a polar compound such as caffeine.



# \* C18 with both silanol activity control and full end-capping is effective in separation of polar compounds.

Sunrise C18 is bonded with octadecylsilane on pure silica gel (average pore size: 12nm, specific surface area: 340m<sup>2</sup>/g), and end-capped after silanol activity control. Final carbon content of Sunrise C18 is 15%.

### ■ Separation of organic acid (Sunrise C18)



Ligand density of Sunrise C18 is intentionally rather low and uniformity of ligands is high, so that it shows enough retention, even if a mobile phase with a low organic solvent content is used, and good peak shape for a polar compound.



5 6

retention time / min

2 3

### Separation of nucleic acid bases (Sunrise C18)

Sunrise C18 and C18-SAC

Silanol Activity Controlled C18 HPLC Column

### Multiple mode separation is achieved on Sunrise series

### \* Silanol groups controlled its activity functions as ion-exchange groups

Sunrise C18-SAC is bonded with octadecylsilane on a pure silica gel and controlled its silanol activity without end-capping. Its carbon content is 14%. Separation on Sunrise C18-SAC is done including hydrogen bond and ion-exchange interaction based on silanol groups except for hydrophobic interaction. Control of pH and salt concentration of a mobile phase can regulate retention.

Separation of basic compounds with ammonium

acetate: Effect of salt concentration(Sunrise C18-SAC)



# ■ Relationship between buffer concentration and retention(Sunrise C18-SAC)

#### Column size: 4.6x150 mm 2 Mobile phase: 70/30=CH<sub>3</sub>CN 3 /acetate buffer (pH4.1) 200mM Flow rate: 1.0 mL/min Temperature: 40 °C 2 Sample 1 3 1 = uracil 2 = toluene 100mM 3 = propranolol4 = nortriptyline 2 1 5 = amitriptyline 3 50mM 2 1 5 3 25mM 2 1 5 3 10mM à 10 12 14 18 24 26 16 20 22 retention time / min

### **Ordering infomation**

Inner diameter	length	Sunrise C18, 5µm	Sunrise C18, 3µm	Sunrise C18-SAC, 5µm	Sunrise C18-SAC, 3µm
[mm]	[mm]	Cat. No.	Cat. No.	Cat. No.	Cat. No.
2.0	50	SB3241	SB2241	SA3241	SA2241
	75	—	SB2251	_	SA2251
	100	SB3261	SB2261	SA3261	SA2261
	150	SB3271	SB2271	SA3271	SA2271
4.6	10	SB3411	SB2411	SA3411	SA2411
	50	SB3441	SB2441	SA3441	SA2441
	75	-	SB2451	-	SA2451
	100	SB3461	SB2461	SA3461	SA2461
	150	SB3471	SB2471	SA3471	SA2471
	250	SB3481	_	SA3481	_
10.0	250	SB3781	_	SA3781	_
20.0	250	SB3881	_	SA3881	_

ChromaNik Technologies Inc.

6-3-1 Namiyoke, Minato-ku, Osaka 552-0001 Japan

TEL: +81-6-6581-0885 FAX: +81-6-6581-0890