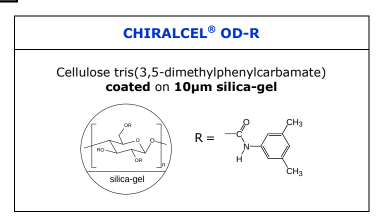




# INSTRUCTION MANUAL FOR CHIRALCEL® OD-R

## Please read this instruction sheet completely before using these columns

## **Column Description**



### Shipping solvent: MeOH

All columns have been pre-tested before packaging. Test parameters and results, as well as the Column Lot Number, are included on a separate (enclosed) page.

#### **CAUTION**

The entire HPLC system including the injector and the injection loop must be flushed with a solvent compatible with the column and its storage solvent prior to connecting. Many of the solvents commonly used in HPLC eluents such as acetone, chloroform, DMF, dimethylsulfoxide, ethyl acetate, methylene chloride and THF may DESTROY the chiral stationary phase if they are present, even in residual quantities, in the system. If an auto-sampler is used, then the solvent employed to flush this unit between injections should also be changed and the relevant solvent lines flushed.

# **Operating Conditions**

	<b>250 x 4.6 mm i.d.</b> Analytical columns
Flow rate direction	As indicated on the column label
Typical Flow rate ${\mathbb O}$	~ 1 ml/min
Pressure limitation	Should be maintained < 300 Bar (4350 psi) for maximum column life Adapt flow rates to column size.
Temperature	0 to 40°C

① The maximum flow rate depends on the mobile phase viscosity (mobile phase composition), and should be adjusted in accordance with the pressure upper's limit (i.e. 300 Bar).

### Please contact Chiral Technologies for further assistance before trying any solvents not mentioned below.

### A - Mobile Phases / For UV and Mass Detections

		ACIDIC (AMPHOTERIC) Compounds	<b>NEUTRAL</b> Compounds	BASIC Compounds <b>9</b>
CHIRALCEL® OD-R	Aqueous solution •	HCOOH aq. pH 2.0	Water	20mM NH₄HCO₃ aq. pH 9.0 adjusted with a basic additive <b>0</b>
	Organic modifier	CH₃CN or MeOH or EtOH or IPA		
	Typical starting conditions §	1		% 0% <b>9</b>

<sup>☞</sup> NOTE 1: If you cannot achieve sufficient resolution, try the complementary mobile phases:

### B - Complementary Mobile Phases / For UV detection

		ACIDIC (AMPHOTERIC) Compounds	<b>NEUTRAL</b> Compounds	BASIC Compounds •
CHIRALCEL® OD-R	Aqueous solution <b>①</b>	$50 \text{mM}$ Phosphate Buffer pH 2.0 OR $H_3 PO_4$ aq. pH 2.0 OR $100 \text{mM}$ KPF $_6$ (or NaPF $_6$ ) aq. pH 2.0 adjusted with $H_3 PO_4$	Water	20mM Borate Buffer pH 9.0  OR  20mM Phosphate Buffer pH 8.0 <b>©</b> OR  100mM KPF <sub>6</sub> (or NaPF <sub>6</sub> ) aq.
	Organic modifier	CH₃CN or MeOH or EtOH or IPA		
	Typical starting conditions <b></b>			0% 0% <b>9</b>

The concentration of all the buffering salt should be less than 500mM.

- Refer to **section C** for preparation of aqueous solutions and choice of basic additives.
- ☐ It is recommended to use CH<sub>3</sub>CN to start the investigation.
  - ☐ The elution power of organic modifiers for these columns is in the descending order of  $CH_3CN > EtOH > MeOH: 50\%CH_3CN \approx 65-70\%EtOH \approx 75-80\%MeOH.$
  - ☐ The use of other organic solvents has not been investigated and could be harmful to the columns.
  - The use of alcohols causes the back pressure to be significantly higher compared to CH<sub>3</sub>CN due to their high viscosity in mixtures with water.
- Retention can be adjusted by changing the proportion of CH<sub>3</sub>CN. Retention may be very sensitive to the amount of CH<sub>3</sub>CN present into the mobile phase.
  - Lowering the column temperature may increase the retention time and the selectivity.
  - □ Increasing the column temperature and decreasing the flow rate may increase the resolution.
- □ To maximize column life the use of a guard column is essential when basic conditions are employed.

- ☐ The use of strong basic conditions (> pH 9) must be avoided, as they are known to damage the silica gel matrix.
- When these columns are used at pH > 7, the temperature should be maintained between 5°C and 25°C for maximum column life.
- High percentages of organic modifier in the mobile phase <u>may precipitate the buffering salt</u> from the solution, and lead to consequent clogging of the column (refer to the table below).

Water / Organic Modifier	Buffer solution / Organic Modifier
90 / 10 to 0 / 100	90 / 10 to 15 / 85

On not use the phosphate buffer for pH > 8. When pH 9 is necessary, use the ammonium bicarbonate solution or borate buffer for maximum column life.

### C - Buffer preparation - Examples

Preparation of pH 2 Phosphate buffer:

**Solution A:** 50mM potassium dihydrogenphosphate

3.40g KH<sub>2</sub>PO<sub>4</sub> / FW 136.09, make up the volume to 500ml with HPLC grade water

**Solution B**: phosphoric acid (H<sub>3</sub>PO<sub>4</sub> 85% by weight) Adjust the pH of solution A to a value of 2.0 using solution B.

> Preparation of pH 2 KPF<sub>6</sub> (NaPF<sub>6</sub>) solution:

**Solution A:** 100mM potassium (sodium) hexafluorophosphate

9.20g KPF<sub>6</sub> / FW 184.06 or 8.40g NaPF<sub>6</sub> / FW 167.95, make up the volume to 500ml with HPLC grade water

**Solution B**: phosphoric acid (H<sub>3</sub>PO<sub>4</sub> 85% by weight)

Adjust the pH of solution A to a value of 2.0 using solution B.

Preparation of pH 9 Ammonium bicarbonate solution:

**Solution A**: 20mM ammonium bicarbonate

0.78g NH<sub>4</sub>HCO<sub>3</sub> / FW 78.05, make up the volume to 500ml with HPLC grade water

**Solution B** Basic additive such as diethylamine (DEA), triethylamine (TEA), ammonia (NH<sub>3</sub>) and so on.

\* DEA tends to give better peak shape than other bases.

Adjust the pH of solution A to a value of 9.0 using solution B.

> Preparation of pH 8 Phosphate buffer:

**Solution A**: 20mM potassium hydrogenophosphate

1.74q of K<sub>2</sub>HPO<sub>4</sub> / FW 174.18, make up the volume to 500ml with HPLC grade water

**Solution B**: 20mM potassium dihydrogenophosphate

1.36g KH<sub>2</sub>PO<sub>4</sub> / FW 136.09, make up the volume to 500ml with HPLC grade water.

Adjust the pH of solution A to a value of 8.0 using solution B.

> Preparation of pH 9 Borate buffer:

**Solution A**: 20mM sodium tetraborate decahydrate

3.81g of  $Na_2B_4O_7.10H_2O$  / FW 381.37, make up the volume to 500ml with HPLC grade water

**Solution B**: 20mM boric acid

 $0.62g\ H_3BO_3\ /\ FW\ 61.83$ , make up the volume to 500ml with HPLC grade water

Adjust the pH of solution A to a value of 9.0 using solution B.

### Column Care / Maintenance

- ☐ The use of a guard column is highly recommended for maximum column life.
- Samples should preferably be dissolved in the mobile phase. The mobile phase and the sample solution should be filtered through a membrane filter of approximately 0.5μm porosity to ensure that there is no precipitate before using.
- □ Before disconnecting the column from the HPLC, any traces of salts should be removed by flushing with a mobile phase that does not contain any salts / buffers, e.q. Water/CH<sub>3</sub>CN 60:40 (v/v).
- ☐ If the column is contaminated with non-eluted components, wash it with a mobile phase that does not contain any salts / buffers then with 100% CH<sub>3</sub>CN for 2 hours at 0.3ml/min. If the non-eluting components are more soluble in methanol, this solvent may be used for the washing step.
- $\ \square$  All salts must be flushed out from the HPLC system and the column before changing to 100% CH $_3$ CN or 100% methanol.
- □ When the column is typically used in reversed-phase mode, use Water/CH<sub>3</sub>CN 60:40 (v/v) to store the column. For other applications the column may be stored in the shipping solvent methanol.

#### **Important Notice**

⇒ STRONGLY BASIC solvent additives or sample solutions MUST BE AVOIDED, because they are likely to damage the silica gel used in these columns.

Operating these columns in accordance with the guidelines outlined here will result in a long column life.

⇒ If you have any questions about the use of these columns, or encounter a problem, contact:

In the USA: <a href="mailto:questions@chiraltech.com">questions@chiraltech.com</a> or call 800-6-CHIRAL In the EU: <a href="mailto:cte@chiral.fr">cte@chiral.fr</a> or call +33 (0)3 88 79 52 00

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