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# Application News

### Gas Chromatography

## **Detector-Switching Analysis Using a Capillary Switching Device**

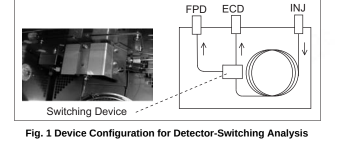
A capillary switching device allows switching of flow lines with high accur acy. This device can be used to switch between 2 detectors, allowing concurrent analyses with m ultiple detectors to be conducted easily. Detector-switching analysis is diff erent from detector-splitting analysis, so when analysis is conducted using detector switching, the entire sample is flowing into the appropriate detector, and accurate information can be obtained from m ultiple detectors during a single analysis r un without sacr ificing sensitivity. (With the detector splitting technique, only a specific fraction of the sample is directed to each of the detectors during the entire analysis.)

#### ■ Analysis of Pesticides Using ECD - FPD Switching

When conducting residual pesticides analysis by GC, a detector with high sensitivity and good selectivity is typically used. Although such a detector is highly effective for analysis of cer tain contaminants in agricultural products, analysis using multiple detectors

A switching program can easily be created with special software that can be do wnloaded from the Shimadzu website free of charge.

No.G266A



is required for detection of all the pesticide constituents. Here we introduce an e xample of sim ultaneous analysis of a standard solution of pesticides using switching between an FPD (flame photometr ic detector) and ECD (electron capture detector).

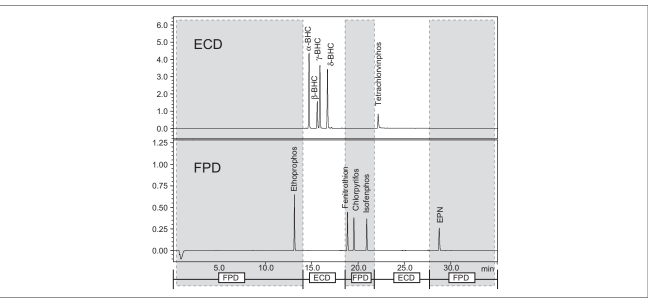


Fig. 2 Chromatograms of Pesticides Obtained by ECD - FPD Switching

**Table 1 Analytical Conditions** 

Instrument	: GC-2010 Plus	Injection Method	: Splitless
Column	: Rtx-5MS (30 m $\times$ 0.25 mmI.D. df = 0.25 $\mu$ m)	Sampling Time	: 1 min (High Pressure: 350 kPa, 1 min)
Column Temp.	: 80 ° C (1 min) - 20 ° C/min - 180 ° C - 5 ° C/min - 280 °	CD@teictor	: ECD: 300 ° C (1nA), Make-up: N2 60 mL/min,
Carrier Gas	: He (150 kPa, Constant Pressure)		FPD: 300 ° C, H2: 80 mL/min, Air: 120 mL/min
Switching Press	s. : 90 kPa	1st Restrictor (ECD side)	) : $0.5 \text{ m} \times 0.18 \text{ mmI.D.}$
Injection Port	: 250 ° C	2nd Restrictor (FPD side	e) : $0.5 \text{ m} \times 0.15 \text{ mmI.D.}$
Sample	: 0.1 mg/L, 2 µL injection		

#### Solvent - Elimination Analysis

Due to the prob lem of diff ering sensitivity among selective detectors, it is advisab le to prevent some solvents or substances from flo wing into a detector, and sometimes these substances cannot be eliminated from the sample . By using a s witching device, unnecessary constituents can be discharged to waste without allowing them to be introduced into the detector. Here we introduce an e xample of analysis in which a solv ent (dichloromethane) is eliminated before entering the detectors; dichloromethane accelerates deterioration of the FTD alkaline source and also has a high response by ECD.

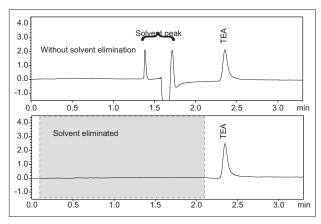


Fig. 3 Chromatograms of With and Without Solvent Elimination

#### Table 2 Analytical Conditions

Instrument	: GC-2010 Plus	Injection Method	: Split, Split Ratio: 1:15
Column	: Rtx-1 (30 m $\times$ 0.32 mmI.D. df = 5 $\mu$ m)	Detector	: FTD: 260 °C 1 pA, H2: 1.5 mL/min, Air: 145 mL/min,
Column Temp.	: 150 °C		Make-up Gas: He27.5 mL/min
Carrier Gas	: He (204.2kPa, Constant Pressure)	1st Restrictor (ECD side)	: $0.5 \text{ m} \times 0.18 \text{ mmI.D.}$
Switching Press.	: 90 kPa	2nd Restrictor (Vent side)	: $0.5 \text{ m} \times 0.15 \text{ mmI.D.}$
Injection Port	: 250 °C		

#### ■ Air Elimination Analysis

The headspace-ECD (HS-ECD) method is used f or analysis of VOCs in water. In this method, both the air that is in vial as w ell as the volatile components are injected. However, since the sensitivity tends to fluctuate when oxygen is introduced into the ECD, this can adversely affect detection stability over time. Using a switching device to discharge air pr ior to reaching the detector can extend the stability of the detector used in the HS-ECD method.

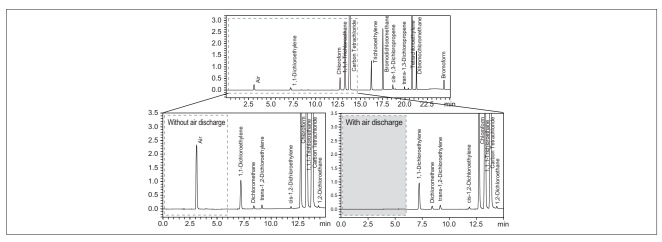


Fig. 4 Chromatograms With and Without Air-Elimination

**Table 3 Analytical Conditions** 

Instrument	: GC-2010 Plus + TurboMatrix HS40	HSVial : 60 °C (60 min)	Injection Time : 0.1 min HS Press : 250kPa
Column	: DB-624 (60 m $\times$ 0.32 mmI.D. df = 1.8 $\mu$ m)	Injection Method : Split	Split Ratio : 1:4
Column Temp.	: 40 °C (5 min) - 4 °C/min - 80 °C - 10 °C/min - 220 °C (3 min)	Detector: ECD 250 1nA	Make-up Gas: (N2) 60mL/min
Carrier Gas	: He (234.4 kPa, Constant Pressure)	1st Restrictor (ECD side)	$0.5m \times 0.18mmI.D.$
Switching Press.	. : 90 kPa	2nd Restrictor (Vent side) :	$0.5 \text{ m} \times 0.15 \text{ mmI.D.}$
Injection Port	: 200 °C		



SHIMADZU CORPORATION. International Marketing Division 3. Kanda-Nishikicho 1-chome, Chiyoda-ku, Tokyo 101-8448, Japan Phone: 81(3)3219-5641 Fax. 81(3)3219-5710 Cable Add.:SHIMADZU TOKYO