### **Environmental Protection Agency**

# Pt. 53, Subpt. B, Table B-1

(C) Both  $P_{20}$  and  $P_{80}$  must be equal to or less than the precision limits speci-

fied in table B–1 to subpart B of part 53 to pass the test for precision.

 $[76\ {\rm FR}\ 54326,\ {\rm Aug.}\ 31,\ 2011,\ as\ amended\ at\ 80\ {\rm FR}\ 65460,\ {\rm Oct.}\ 26,\ 2015]$ 



FIGURE B-1 TO SUBPART B OF PART 53-EXAMPLE

Figure B-1 to Subpart B of Part 53--Example showing the nature of the tracing obtained during the test sequence for 24-hour drift, lag time, rise time, fall time, and precision. The time scale has been greatly compressed.

Performance parameter	Units <sup>1</sup>	SO <sub>2</sub>		O <sub>3</sub>		со		NO <sub>2</sub>	Definitions
		Std. range <sup>3</sup>	Lower range 2 3	Std. range <sup>3</sup>	Lower range 2 3	Std. range <sup>3</sup>	Lower range 2 3	(Std. range)	and test procedures
1. Range 2. Noise 3. Lower detectable limit. 4. Interference equiv-	ppm ppm ppm	0–0.5 0.001 0.002	<0.5 0.0005 0.001	0–0.5 0.0025 0.005	<0.5 0.001 0.002	0–50 0.2 0.4	<50 0.1 0.2	0–0.5 0.005 0.010	Sec. 53.23(a) Sec. 53.23(b) Sec. 53.23(c)

TABLE B–1 to Subpart B of Part 53—Performance Limit Specifications for Automated Methods

limit.									
4. Interference equiv-									
alent									
Each interferent	ppm	±0.005	<sup>4</sup> ±0.005	±0.005	±0.005	±1.0	±0.5	±0.02	Sec. 53.23(d
Total, all	ppm	-	-	-	-	-	-	0.04	Sec. 53.23(d
interferents.									
5. Zero drift, 12 and	ppm	±0.004	±0.002	±0.004	±0.002	±0.5	±0.3	±0.02	Sec. 53.23(e
24 hour.									
6. Span drift, 24 hour									
20% of upper	Percent	-	-	-	-	-	-	±20.0	Sec. 53.23(e)
range limit.									
80% of upper	Percent	±3.0	±3.0	±3.0	±3.0	±2.0	±2.0	±5.0	Sec. 53.23(e
range limit.									
7. Lag time	Minutes	2	2	2	2	2.0	2.0	20	Sec. 53.23(e)
8. Rise time	Minutes	2	2	2	2	2.0	2.0	15	Sec. 53.23(e
9. Fall time	Minutes	2	2	2	2	2.0	2.0	15	Sec. 53.23(e
10. Precision									

#### Pt. 53, Subpt. B, Table B-2

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Performance parameter	Units <sup>1</sup>	SO <sub>2</sub>		O <sub>3</sub>		со		NO <sub>2</sub>	Definitions
		Std. range <sup>3</sup>	Lower range <sup>23</sup>	Std. range <sup>3</sup>	Lower range <sup>23</sup>	Std. range <sup>3</sup>	Lower range 2 3	(Std. range)	and test procedures
20% of upper range limit.		-	-	-	-	-	-		Sec. 53.23(e)
-	Percent <sup>5</sup>	2	2	2	2	1.0	1.0	4	Sec. 53.23(e)
80% of upper range limit.		-	-	-	-	-	-		Sec. 53.23(e)
-	Percent <sup>5</sup>	2	2	2	2	1.0	1.0	6	Sec. 53.23(e) Sec. 53.23(e)

<sup>1</sup>To convert from parts per million (ppm) to ug/m<sup>3</sup> at 25 °C and 760 mm Hg, multiply by M/0.02447, where M is the molecular weight of the gas. Percent means percent of the upper measurement range limit. <sup>2</sup>Tests for interference equivalent and lag time do not need to be repeated for any lower range provided the test for the standard range shows that the lower range specification (if applicable) is met for each of these test parameters. <sup>3</sup>For candidate analyzers having automatic or adaptive time constants or smoothing filters, describe their functional nature, and describe and conduct suitable tests to demonstrate their function aspects and verify that performances for calibration, noise, lag, rise, fall times, and precision are within specifications under all applicable conditions. For candidate analyzers which operator-selectable time constants or smoothing filters, conduct calibration, noise, lag, rise, fall times, and precision the FRM designation. <sup>4</sup>For nitric oxide interference for the SO<sub>2</sub> UVF method, interference equivalent is ±0.0003 ppm for the lower range. <sup>5</sup>Standard deviation expressed as percent of the URL.

### [80 FR 65460, Oct. 26, 2015]

TADT D 0		T D OT DAD		
TABLE B-Z	TO SUBPAR	г в ог рак	$T_{00} = T_{EST}$	ATMOSPHERES

Test gas	Generation	Verification			
Ammonia	Permeation device. Similar to system described in references 1 and 2.	Indophenol method, reference 3.			
Carbon dioxide	Cylinder of zero air or nitrogen containing CO <sub>2</sub> as required to obtain the concentration specified in table B-3.	Use NIST-certified standards whenever possible. If NIST standards are not available, obtain 2 stand- ards from independent sources which agree within 2 percent, or obtain one standard and submit it to an independent laboratory for analysis, which must agree within 2 percent of the supplier's nominal analysis.			
Carbon monoxide	Cylinder of zero air or nitrogen containing CO as re- quired to obtain the concentration specified in table B–3.	Use an FRM CO analyzer as described in reference 8.			
Ethane	Cylinder of zero air or nitrogen containing ethane as required to obtain the concentration specified in table B–3.	Gas chromatography, ASTM D2820, reference 10. Use NIST-traceable gaseous methane or propane standards for calibration.			
Ethylene	Cylinder of pre-purified nitrogen containing ethylene as required to obtain the concentration specified in table B–3.	Do.			
Hydrogen chloride	Cylinder <sup>1</sup> of pre-purified nitrogen containing approxi- mately 100 ppm of gaseous HCI. Dilute with zero air to concentration specified in table B–3.	Collect samples in bubbler containing distilled water and analyze by the mercuric thiocyanate method, ASTM (D612), p. 29, reference 4.			
Hydrogen sulfide	Permeation device system described in references 1 and 2.	Tentative method of analysis for H <sub>2</sub> S content of the atmosphere, p. 426, reference 5.			
Methane	Cylinder of zero air containing methane as required to obtain the concentration specified in table B-3.	Gas chromatography ASTM D2820, reference 10. Use NIST-traceable methane standards for cali- bration.			
Nitric oxide	Cylinder <sup>1</sup> of pre-purified nitrogen containing approxi- mately 100 ppm NO. Dilute with zero air to re- quired concentration.	Gas phase titration as described in reference 6, section 7.1.			
Nitrogen dioxide	<ol> <li>Gas phase titration as described in reference 6</li> <li>Permeation device, similar to system described in reference 6.</li> </ol>	<ol> <li>Use an FRM NO<sub>2</sub> analyzer calibrated with a grav- imetrically calibrated permeation device.</li> <li>Use an FRM NO<sub>2</sub> analyzer calibrated by gas- phase titration as described in reference 6.</li> </ol>			
Ozone	Calibrated ozone generator as described in reference 9.	Use an FEM ozone analyzer calibrated as described in reference 9.			
Sulfur dioxide	<ol> <li>Permeation device as described in references 1 and 2.</li> <li>Dynamic dilution of a cylinder containing approxi-</li> </ol>	Use an SO <sub>2</sub> FRM or FEM analyzer as described in reference 7.			
Water	mately 100 ppm SO <sub>2</sub> as described in Reference 7. Pass zero air through distilled water at a fixed known temperature between 20° and 30 °C such that the air stream becomes saturated. Dilute with zero air to concentration specified in table B–3.	Measure relative humidity by means of a dew-point indicator, calibrated electrolytic or piezo electric hygrometer, or wet/dry bulb thermometer.			