Indian Standard

METHODS OF PHYSICAL TESTS FOR HYDRAULIC CEMENT

PART 9 DETERMINATION OF HEAT OF HYDRATION

(First Revision)

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0. FOREWORD

- 0.1 This Indian Standard (Part 9) (First Revision) was adopted by the Bureau of Indian Standards on 22 April 1988, after the draft finalized by the Cement and Concrete Sectional Committee had been approved by the Civil Engineering Division Council.
- **0.2** Standard methods of testing cement are essential adjunct to the cement specifications. This standard in different parts lays down the procedure for the tests to evaluate the physical properties of different types of hydraulic cements. The procedure for conducting chemical tests of hydraulic cement is covered in IS: 4032-1985*.
- 0.3 Originally all the tests to evaluate the physical properties of hydraulic cements were covered in one standard; but for facilitating the use of this standard and future revisions, it has been decided to print the different tests as different parts of the standard and accordingly, this revised standard has been brought out in thirteen parts. This will also facilitate updating of individual tests. Further,

since publication of the original standard in 1968 a number of standards covering the requirement, of different equipment used for testing of cement, a brief description of which was also covered in the standard, had been published. In this revision, therefore, reference is given to different instrument specifications deleting the description of the instruments, as it has been recognized that reproducible and repeatable tests results can be obtained only with standard testing equipment capable of giving desired level of accuracy. This part (Part 9) covers determination of heat of hydration of cement.

0.4 For the purpose of deciding whether a particular requirement of this standard is complied with, the final value, observed or calculated, expressing the result of a test or analysis, shall be rounded off in accordance with IS: 2-1960*. The number of significant places retained in the rounded off value should be the same as that of the specified value in this standard.

1. SCOPE

1.1 This standard (Part 9) covers the procedure for determining the heat of hydration of cement as expressed in kilojoules per kilogram.

2. SAMPLING AND SELECTION OF TEST SPECIMENS

2.1 The samples of the cement shall be taken in accordance with the requirements of IS: 3535-1986* and the relevant standard specification for the type of cement being tested. The representative sample of the cement selected as above shall be thoroughly mixed before testing.

3. TEMPERATURE

3.1 The temperature of moulding room, dry materials, appliances and water shall be maintained at $27 \pm 2^{\circ}$ C.

4. APPARATUS

- **4.1 Calorimeter** Calorimeter conforming to IS: 11262-1985*.
- 4.2 Mortar and Pestle Approximately 200 mm in diameter mortar and pestle for grinding partially hydrated samples.
- 4.3 Glass/Plastic Vials Glass/plastic vials having the dimension approximately 80×20 mm with tight fitting stoppers or caps.

^{*}Method of chemical analysis of hydraulic cement (first revision).

^{*}Rules for rounding off numerical values (revised).

[•]Methods of sampling hydraulic cements (first revision).

^{*}Specification for calorimeter for determination of heat of hydration of hydraulic cement.

- 4.4 Stop Watch or Timer The timer shall have a positive starting and stopping mechanism and shall be capable of being read to the nearest 0.5 s or less. The timer shall be accurate to 0.5 s or less for time interval up to 60 s and to 1 percent or less for time intervals of 60 to 300 s.
- **4.5 Sieve** 150 μ m and 850 μ m IS sieve conforming to IS: 460 (Part 1) 1985*.
- **4.6** Muffle Furnace Muffle furnace capable of maintaining a temperature of 900 to 950°C.
- **4.7** Analytical Balance Analytical balance capable of reproducing results within 0.0002 g with an accuracy of ± 0.0002 g.

Note — Self-indicating balance with equivalent accuracy may also be used.

4.8 Standard Weights

4.9 Weighing Bottles

4.10 Camel Hair Brush

5. MATERIAL

- 5.1 Nitric Acid of 200 ± 0.05 N strength, made in bulk from analytical reagent quality materials. Whenever a new batch is prepared, the heat capacity of the calorimeter shall be redetermined.
- 5.2 Hydrofluoric Acid 40 percent (w/w), analytical reagent quality.
- 5.3 Zinc Oxide Analytical reagent quality.
- 5.4 Wax paraffin wax.
- 5.5 Distilled Water conforming to IS: 1070-1977†.

6. PROCEDURE

6.1 Determination of the Heat Capacity

- 6.1.1 Inspect the wax lining for faults. Measure into the calorimeter 9.6 ± 0.1 ml of hydrofluoric acid and 388.0 ± 0.1 ml of 2.0 N nitric acid at a temperature of $27 \pm 2^{\circ}$ C. For convenience in measuring the nitric acid, a special standard flask of 388 ml capacity calibrated at 27° C shall be constructed. For measuring the hydrofluoric acid, a small measuring cylinder shall be made up by sealing a 15-cm length of 1-cm diameter 'polythene' resin tube to a flat plate of the same material with a small gas jet.
- 6.1.2 Take a quantity of zinc oxide sufficient for about six determinations. Ignite it for one hour at 900 to 950°C, cool in a desiccator con-

taining anhydrous calcium chloride and grind it to pass a 150 micron IS Sieve. For each determination, about 7.0 g of this ignited oxide shall again be heated to 900 to 950°C for 5 min and then cooled for not less than $2\frac{1}{2}$ h and not more than 5 h in the desiccator containing anhydrous calcium chloride before weighing accurately.

- 6.1.3 Assemble the calorimeter and run the stirrer for at least 5 min to allow the temperature to become uniform. Take temperature reading correct to 0.001°C every minute for 5 min to determine the initial heating or cooling correction. Then introduce the zinc oxide from the funnel steadily over a period of 1 to 2 min. The funnel shall then be brushed clean with camelhair brush. Take temperature readings at one minute intervals until the solution is complete, as indicated by a steady rate of heating of cooling of the calorimeter. The solution period shall not exceed 20 min. Continue the readings for a further period of 5 min, to determine the final heating or cooling correction.
- 6.1.4 Plot initial and final heating or cooling rates against the corresponding calorimeter temperature, namely the Beckmann readings at the beginning of the solution period and at the end, respectively. Join the two points by a straight line (see Fig. 1 and example in 7.1). From this graph, the corrections are read off for each temperature reading during the solution period. These corrections shall be summed and the total added or subtracted as appropriate to the observed temperature-rise.

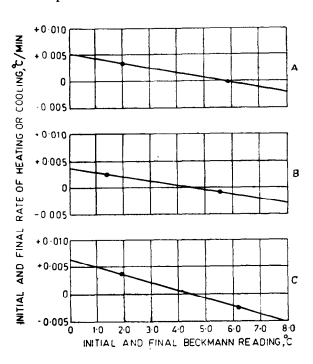


Fig. 1 Heating or Cooling Corrections

^{*}Specification for test sieves: Part 1 Wire cloth test sieves (third revision).

[†]Specification for water for general laboratory use (second revision).

6.1.5 Calculate the heat capacity as follows: Heat capacity

$$(J/^{\circ}C) = \frac{\text{Mass of ZnO (g)}}{\text{Corrected temperature-rise}} \times [1 072 + 0.4 (30 - \phi) + 0.5 (\phi_{\circ} - \phi)]$$

where

1 072= heat of solution of zinc oxide at 30°C,

0.4 = negative temperature coefficient of the heat of solution,

 final temperature of the calorimeter and contents in °C.

0.5 = specific heat of zinc oxide, and

 ϕ_{\circ} = room temperature in °C.

This expression simplifies to:

Heat capacity

$$= \frac{\text{Mass of ZnO (1 084 - 0.9 } \phi + 0.5\phi_{\circ})}{\text{Corrected temperature-rise}}$$

6.2 Preparation of Cement Sample — Mix by hand for 4 min, 60 g of cement and 24 ml of distilled water which shall be between 15 and 25° C. Fill with this mixture 3 glass/plastic vials, cork and then seal with wax. Store the specimen vials with the mixture in a vertical position at $27 \pm 2^{\circ}$ C.

6.3 Determination of the Heat of Solution

6.3.1 For determination of the heat of solution of unhydrated cement, weigh a sample of about 3.0 g. At the same time, weigh out another quantity approximately 7.0 g for the loss on ignition. Both the weighings shall be correct to the nearest 0.001 g. Carry out the determination of temperature-rise exactly as described for zinc oxide.

Calculate the heat of solution from the following formula:

Heat of solution (kJ/kg) of unhydrated cement

where 0.8 is the specific heat of unhydrated cement.

- **6.3.2** The mean of three determinations which shall be carried out within 7 days of the mixing of the hydrated samples shall be taken.
- 6.3.3 For the determination of heat of solution of hydrated cement, break open one of the glass vials (see 6.2). Remove the adherent wax and glass from the cement, then grind the cement (as rapidly as possible to avoid carbonation) to pass an 850-micron IS Sieve. Keep the ground sample in a stoppered weighing bottle from which weigh out samples of 4.2 and 7.0 g for heat of solution and loss on ignition, respectively. The

loss on ignition shall be determined on each sample used for heat of solution. Carry out the determination of temperature-rise as before and calculate the heat of solution from the following formula:

Heat of solution of hydrated cement (kJ/kg ignited mass)

Heat capacity × corrected temperature-rise

Mass of sample corrected for ignition loss

-1.7 (
$$\phi_0 - \phi$$
)

where 1.7 is the specific heat of hydrated cement.

The mean of three determinations on separate vials shall be taken.

6.4 Ignition Loss — Place the sample in a cool furnace and raise the temperature of the furnace to 900° C over a period of one hour. Keep the sample at $900 \pm 50^{\circ}$ C for 3 to 4 h and then cool it in a desiccator containing anhydrous calcium chloride. Weigh after half an hour. All weighings shall be correct to the nearest milligram.

7. CALCULATION

7.1 Calculate the heat of hydration by subtracting the respective heats of solution of hydrated cement from the heat of solution of the unhydrated cement. The heats of hydration shall be determined at 7 and 28 days. Heats of solution shall be calculated to the nearest 0.5 kJ/kg and heats of hydration to the nearest 5 kJ/kg as given in the following example:

Example:

a) Determination of heat capacity

Time	°C Beckmann Calorimeter	Heating or Cooling Correction
(min)	Temperature	(see Graph C, Fig. 1)
0 1 2 3 4 5	1.891 1.894 1.898 1.902 1.905 1.908	Initial correction +0.003 4
6 7 8 9 10	2:550 5:880 6:175 6:225 6:241 6:245	$ \begin{array}{r} -0.002 \ 4 \\ +0.002 \ 2 \\ +0.002 \ 5 \\ +0.002 \ 5 \\ +0.002 \ 6 \\ 0.002 \ 6 \\ = +0.010 \ 0 \end{array} $
12 13 14 15 16	6.243 6.240 6.237 6.234 6.232 6.230	Final correction -0.002 6
Temperat	ure-rise = 6°2	245 - 1.908 = 4.337

Temperature-rise = 6.245 - 1.908 = 4.337Correction = + 0.010

IS: 4031 (Part 9) - 1988

Corrected temperature-rise = 4.347

Mass of zinc sample = 7.00 g

Room temperature = 27.00°C

Final temperature of calorimeter and contents* = 27.75°C

Heat capacity = $\frac{7.00}{4.347} \times$ (1.084 - 0.9 × 27.75 + 0.5 × 27) = $\frac{7.00}{4.347} \times$ (1.084 - 24.975 + 13.5) = $\frac{7.00}{4.347} \times$ 1.072.525 = 1.727 J/°C

b) Determination of heat of solution on anhydrous cement sample

en:	0.5 D I			= 2
Time	°C Beckmann	Heating or Cooling	a) D a	etermination of
(:)	Calorimeter	Correction		ated cement
(min)	Temperature	(see Graph B, Fig. 1)		orage at 27°C
0	1.552		310	rago at 27 C
1	1.528		Time	°C Beckman
2	1:230	Initial correction	2	Calorimeter
2 3 4	1.232 f 1.234 f	+0.0022	(min)	Temperatur
5	1.234			
	1 230)		0	2.019
6	3.350	-0 000 6	1	2.022
7	4.460	+0.0005	2 3	2:026 L
8	4.850	+0.000 5	3 4	2.030
9	5.090	$+0.000\ 6$	5	2 035
10	5.230	+0.000 7	3	2 033)
11 12	5:330	+0.000 7	6	5.000
12	5·392 5·432	+0.000 7	7	5·700
13	5·460	+0.000 7 +0.000 7		
15	5· 4 75	+0.000 8	8	5.815
16	5.483	+0.000 8	9	5.845
17	5 489	+0.000 8	10	5.828
18	5 491	+0.000 8	11	5.867
19	5.492	+0.0008	12	5.872
		$\frac{+0.008}{}$	13	5.877
•0	5,4003	,	14	5.880
20	5:492		15	5.881
21 22	5·491 5·490	Final correction	16	5.882
23	5·490 }	-0.000 8	10	3 002
24	5.490	0 000 0		
25	5.488		17	5.882
26	5·487 j		18	5·882 İ
	_		19	5.882
Temperatu		92 - 1.236 = 4.256	20	5.885 >
Correction	ı = +	0.008	21	5.882
Corrected			22	5.881
tempera	ture-rise = 4·2	64	23	ر 5٬880

^{*}Determined separately by a mercury-in-glass thermometer.

Mass of cement = 3.000 gsample = 1.91 percent Ignition loss Room temperature = 27.00°C Final temperature of calorimeter and contents* $= 27.25^{\circ}C$ Heat capacity of $= 1^{\tilde{7}27} \text{ J/°C}$ calorimeter anhydrous cement = $\frac{1.727 \times 4.264 \times 100}{2}$ Heat of solution of -0.8(27.00 - 27.25) $\frac{1.727 \times 4.264 \times 100}{20.20} + 0.2$ 3.000×98.09 = 2502.4 + 0.2= 2 502.6 kJ/kg

 c) Determination of heat of solution on hydrated cement sample after 28 days' storage at 27°C

= 2 502.5 kJ/kg (say)

sto	orage at 27°C			
Time	°C Beckmann Calorimeter	Heating or Cooling Correction		
(min)	Temperature	(see Graph A, Fig. 1)		
0 1 2 3 4 5	2·019 2·022 2·026 2·030 2·032 2·035	Initial correction +0.003 2		
6	5.000	-0·000 5		
7	5.700	+0.000 2		
8	5.815	+0.000 5		
ğ	5.845	+0.0002		
10	5.828	+0.000 2		
11	5.867	+0.000 2		
12	5.872	+0.000 3		
13	5.877	+0.000 3		
14	5.880	+0.000 3		
15	5.881	+0.000 3		
16	5.882	+0.000 3		
		+0.002		
17 18 19 20 21 22	5.882 5.882 5.882 5.882 5.882 5.881	Final correction -0.000 3		
23	5 [.] 880 J			

^{*}Determined separately by a mercury-in-glass thermometer.

Temperature-rise = 5.882 - 2.035 = 3.847Heat of solution of $=\frac{1.727 \times 3.849 \times 100}{1.000}$ hydrated cement = 0.002Correction 4.200 0 × 72 04 Corrected -1.7(27.00 - 24.5)temperature-rise = 3.849 $=\frac{1.727 \times 3.849 \times 100}{4.200}$ Mass of cement $4.200 \ 0 \times 72.04$ = 4.2000 gsample Ignition loss = 27.96 percent -425= 2 196.93 - 4.25Room temperature* = 27 00°C = 2 192.68 kJ/kgFinal temperature of calorimeter and = 2 192 5 kJ/kg (say)= 24.5°C contents*

Heat capacity of calorimeter

= 1.727 J/°C

d) Determination of heat of hydration

Heat of hydration at

= 2502.5 - 2192.528 days = 310.0 kJ/kg

^{*}Determined separately by a mercury-in-glass thermometer.

Bureau of Indian Standards

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