For qualitative estimation of the Erythrocyte Sedimentation Rate (ESR or Sed-Rate).

For In Vitro Diagnostic Use Only

Contents: (sufficient for 100 ESR determinations)
100 Dispettes* (Westergen dimensions) plus 100 blue filling reservoirs.

*Note: The word Dispette or Dispettes used throughout this document is the Trade Name for the Disposable ESR Pipette manufactured by Guest Scientific AG.

Storage Conditions:
The reservoirs and Dispettes may be stored at room temperature, which may be cold, cool or warm: However the test itself must be performed at temperatures between 18 – 25°C (64 – 77°F) see note c) on page 3.

Background: The erythrocyte sedimentation rate (ESR) is one of the most widely requested laboratory tests throughout the world and manual methods, such as the Dispette, are particularly popular in the outpatient’s setting in the USA. While infection may be the most common cause of elevated results, many other conditions such as malignant tumors and renal disease have been associated with raised values; hence its role has become established in the Clinician’s mind as a screening test for the presence of clinical illness. Elevated ESR results correlate well to the severity of acute inflammatory disease and the test has been cited as a useful indicator of stroke severity and predictor of early relapse or survival in Hodgkin Disease.

In 1926 Alf Westergren, published ‘The Technique of the Red Cell Sedimentation Reaction’, based on the work of Robin Fahræus in 1918. His technique included the use of a combined diluted and anticoagulant (tri-sodium citrate) for the blood sample and specified the length (200mm) and bore diameter (2.5mm) of the measuring tube. His technique became the basis of the test we know today as the Westergen method and the pipet dimensions and dilution factor were adopted as the reference method for erythrocyte sedimentation rate by the International Committee for Standardization in Hematology (ICSH) in 1973.

Since 1926 various workers have proposed variations to the Westergen method, (e.g. Wintrobe in 1935 and the ‘zeta’ sedimentation rate in 1972), however in 1993 the ICSH stated that a slightly modified Westergen method should be used for the reference method to which all other techniques, if not conforming, should be standardised. The most recent review by the ICSH published in April 2011 confirms that the reference method for measurement of the ESR should be based on the Westergen method using diluted blood. In similar manner the most recent CLSI Approved Standard (5th Ed.), for ESR recognises the Westergen method as the standardized or selected procedure, making it a truly ‘International’ standard.

The dimensions of the Dispette conform to those of the Westergen method.

Physical basis of blood sedimentation: To this day the phenomenon of erythrocyte sedimentation is still only partly understood, however three definite phases of the process have been identified: During the first, or Lag Phase, the red cells form a characteristic rouleaux pattern and sedimentation is generally slow. The rate accelerates in the second period, the Decantation Phase and slows again in the final Packing Phase as the red cell aggregates pile up towards the lower part of the tube.

Note: Sedimentation is not linear and the time taken over each of the Lag, Decantation and Packing phases will differ between patients, hence the observer must never try to ‘estimate’ the final result before the full 60 minute time period for the test has elapsed. The size of rouleaux aggregates formed in the Lag Phase is the critical factor affecting the final result. Rouleaux appears to be mainly influenced by plasma proteins including fibrinogen, IgM and alpha2-macroglobulin. Opinions vary as to the accelerating and retarding properties of glycoprotein and albumin. IgG appears to increase the sedimentation rate only at high concentrations.

Items required but not provided in the Dispette box:
1. Transfer pipettes.
2. Sodium citrate (see below) or Normal saline.
3. An ESR stand to hold the Dispettes and reservoir assemblies in a vertical position, controlled by means of a levelling bubble and adjusting screws (code: GS or FH 1705).
4. A laboratory or similar timer capable of alarming after exactly one hour (60 minutes) elapsed time.

Sample requirement:
Whole blood should be obtained by clean venipuncture over a maximum period of 30 seconds without excessive stasis. A traditional syringe or a vacuum extraction system may be used. The blood sample may be taken into an EDTA tube or a pre-citrate sample tube (using 3.3% - 0.109 mol/L, tri-sodium citrate; CaH2O4Na2*2 H2O; CAS number 6132-04-3) to give a dilution ratio of 1 part citrate to which is added 4 parts blood sample. The blood sample and anticoagulant mixture must be immediately mixed gently, but thoroughly at least 8 times by complete inversion of the sample tube.

The most recent ICSH review states that citrated blood samples must be tested within 2 hours if left at room temperature, or 4 hours if stored at 4°C. Note: In the USA current CLSI guidance states that the blood sample may be stored for up to 24 hours in a refrigerator. Blood specimens stored in a refrigerator must return to room temperature naturally before testing starts.

If the sample is left for any amount of time before testing, either at room or refrigerator temperature, it must be well mixed by at least 8 gentle inversions of the tube before transfer to the reservoir.
Instructions for Use:

**Note:** Always wear appropriate personal protective clothing as indicated by Good Laboratory Practice. If blood is spilled follow your laboratory policy or national guidance, for safe disposal.

1. Mix the blood sample thoroughly but gently, by at least 8 full inversions of the container.

2. **Pre-citrated blood sample:** using a transfer pipette add 1.25ml of the well mixed sample into the filling reservoir. Ensure the sample volume does not go beyond the filling line (diagram 1).

3. **EDTA blood sample:** In a 12 x 75mm test tube add 4 parts of the blood sample to 1 part Normal saline solution (0.85%-0.145 mol/L, NaCl). Mix very thoroughly, but carefully, by at least 8 full inversions of the test tube. Transfer 1.25ml of this mixture to the blue filling reservoir – ensure that the volume of sample reaches but does not go beyond the filling line (diagram 1).

4. Without delay gently insert the Dispette into the reservoir and observe the blood sample being pushed up inside the bore until it reaches to or beyond the zero level (see diagrams 2 & 3). The Dispette must be fully inserted to the bottom of the reservoir – if the blood level in the Dispette then reaches above the zero mark on the graduated scale, apply gentle pressure with the thumb to the side of the tube (see diagram 4) and the blood level will drop – stop at exactly the zero mark.

5. Place the full Dispette assembly (Dispette plus reservoir) in a correctly levelled ESR stand ensuring that the Dispette is at 90 degrees, plus or minus no more than one degree to the horizontal, (i.e. vertical, see diagrams 5a & 5b). **Immediately start the timer.**

6. At the end of the timed hour, the result is read by aligning the eye to the level where the red cell column has dropped – leaving clear plasma above (see illustration d on page 3). Record the number of mm the red cells have dropped from the mm scale printed on the Dispette.

7. Record the result as ESR = $x$ mm
   
   or Sed-Rate = $x$ mm

**Note:** It is common, though an incorrect practice, to record ESR results as $x$ mm/hour or $x$ mm per hour. Both the ICSH and CLSI point out that despite the common title Erythrocyte Sedimentation Rate, it is not in fact a rate but a measurement at one hour. Hence both these official organizations currently recommend that results should be reported as: ESR = $x$ mm.
PRECAUTIONS:

a) Thorough mixing of the blood sample before adding to the reservoir is essential. However it must be done gently, shaking the sample will result in haemolysis which may obscure the end point.

b) Shaking the sample may also create bubbles which will seriously affect the result. If bubbles are present in the Dispette repeat the test with a fresh sample that has been more carefully mixed.

c) The ESR must be performed at room temperature (defined by the ICSH as 18 to 25°C or 64 to 77°F). Do not place the stand near a window, in direct sunlight or where it may be subjected to drafts.

   Note: Should it be impossible to perform the test within this temperature range request a temperature correction chart from Arkray USA Inc.

d) The ESR is affected by vibration; ensure the stand is placed well away from machinery and that the bench is not subjected to knocks. Remember that vibration may only occur intermittently and/or may be as a result of machinery (e.g. a centrifuge) sited in another part of the building.

e) Do not pick up the stand to read the result as this will affect other tests in progress. Bring the eye to the level of the red cells to read accurately from the scale (see illustration d).

f) If the result falls between 130 and 155mm, the exact end point may be obscured by the filling reservoir. In this case report the result as >130 mm. If a numerical value is requested estimations can be made, by reading from the scale of an unused Dispette pipette held next to the test assembly.

   Note: Before reporting an estimated result be sure to verify that this complies with your laboratory's best practice and standards. Record and report that the result has been estimated.

g) Occasionally the level of the red cells is not clear-cut and a ‘Christmas Tree’ effect may be observed. In such cases the level where the red cells become fully concentrated should be recorded (see illustration e).

h) In cases of serious infection or leukemia a heavy layer of white cells may be present on top of the column of red cells. This should be ignored and the reading taken from the level of the red cells only.

i) It must be remembered that the red cells will continue to settle after one hour has elapsed. Therefore it is most important to read the test at exactly one hour after setting up the test in the stand (e.g. the test reading will generally be higher at 1 hour 15 minutes than at 1 hour).

j) Note: The rate of fall of the red cells is not linear. Therefore it must not be assumed that the reading at 60 minutes will be twice that at 30 minutes (see Physical basis of blood sedimentation on page 1).
Calibration, Quality Control and Normal Values:

For calibration we recommend reference to the ICSH Review of Measurement of erythrocyte sedimentation rate, published in April 2011,¹¹ the original ICSH paper published in 1993,¹⁰ or the U.S. equivalent: CLSI guidance document H02-A5.¹²

Some commercial companies offer quality control samples for ESR; however we have not been able to compare them to the ICSH reference method. It is recommended that laboratories must therefore instigate and record their own quality control system in accordance with ICSH or CLSI guidance and/or their own country’s regulatory requirements.

As normal values for the ESR can vary with age and ethnic origin, reference to the ICSH or CLSI document should also be made for guidance on normal values appropriate to individual countries.

For a list of Frequently Asked Questions about ESRs please go to www.guestscientific.com & follow the links from the drop down menu.

Text References