# TECHNICAL PAPER

# JOURNAL OF THE SOUTH AFRICAN INSTITUTION OF CIVIL ENGINEERING

Vol 58 No 2, June 2016, Pages 14—24, Paper 1232



PHLIP STOTT (Pr Eng. MSNCE) received BSc. (Hons) and MSc degrees from Manchester University. He lectured at Ahmadu Bello University in Nigeria and the University of the Witwatersyand in Johannesburg, and has been working as a consulting engineer since 1984. He received the Henry Adams Award from the Institute of Structural Engineers. London.

Currently he is a DTech Candidate and a member of the Soil Mechanics Research Group at the Central University of Technology (CUT), Bloemfontein, He is also a member of the Structural and Geotechnical Divisions of SAICE (South African Institution of Civil Engineering).

Contact detaik:
Department of Cwl Engineering
Central University of Technology
Phylate Bag X2039
Bloemfortish 9300
South Africa
T: +27 82 253 4001
E: philip@f10.co.za



DR ELIZABETH THERON (Pr Tech Eng) received an NHDjp and MTech in Civil Engineering from the then Technikon Free State (renamed since to the Central University of Technology, or CUT for short). Bloemforkein, South Africa. She also has a PhD in Geography from the University of the Free State, Bloemfontein. She has lectured in civil engineering at the forerunner of the CUT

civil engineering at the forerunner of the CUT since 1987, and has received the Prestige Award of the Vice-Chancellor (Academic; Class Gold). She is currently a senior lecturer at the CUT, and project manager of, and researcher in, CUT's Soll Mechanics Research Group project manager of, and researcher in, CUT's Soll Mechanics Research Group

Contact delais:
Department of Civil Engineeling
Central University of Technology
Privane Bag X20539
Bloernfornlein 9900
South Africa
T: 427 \$1 \$07 3646
E: etheron@cut.ac.za

Keywards: hydrometer analysis, clay fraction, dispersion of clays, de-flocculation

#### Note to readers:

This paper contains 18 supporting photographs which are discussed in detail on pages 20–22. However, due to space constraints, the photographs have been spread evenly throughout the paper. We trust that this will not inconvenience our readers.

# Shortcomings in the estimation of clay fraction by hydrometer

P Statt, E Theror

with and without labelling of clay minerals using an exchangeable cation dye specified hydrometer procedures. Clays are examined both with and without dispersant, and adequacy of dispersion and possible consequences for clay fraction determination in currently deals with a microscopic examination of some typical South African clayey soils to assess the the standard procedures for assessment of clay fraction in well over half a century. This paper the possible sources of error have been suggested, but little or no change has been made in a hydrometer, It has long been suspected that there may be flaws in this approach. Some of soil. SANS 3001 GR3 (SANS 2011) specifies a procedure for clay fraction determination using The estimation of clay fraction is important for predicting the engineering properties of a

#### INTRODUCTION

four factors (Savage 2007): in its efficiency. Savage suggested that the hydrometer method may be doubtful due to countries. It is, however, somewhat dubious Britain, America, Australia and many other 2011), is very similar to that specified in by hydrometer, as specified in the South African standard SANS 3001 GR3 (SANS The method of estimating clay fraction defined as PI/clay fraction (Skempton 1953) and clay fraction. Skempton's "activity" is Merwe 1964) uses the plasticity index (PI) design. Van der Merwe's method (Van der heave potential relating to foundation including common methods of assessing is required for a number of soil evaluations, An estimation of the clay fraction of a soil

- Stoke's law assumes all particles to be spherical, while clays are flaky.
- 2. De-flocculation of many clays is seldom
- fully completed at the time of testing.

  3. Clay particles are partially carried down

by the larger particles.

have found wide acceptance error may be. The method does not appear to better estimate, or what the likely margins of appears to be no way of telling which gives the er, some lower than the hydrometer). There clear pattern of correlation (some values high-Soil Mechanics Research Group revealed no by the Central University of Technology (CUT) give examples, and the examination of samples ter results and Savage's method. Savage did not clear pattern of correlation between hydromeformula. Unfortunately there seems to be no fraction indirectly by using Skempton's activity Savage proposed a method of estimating clay ₽ all particles, which may not be true. A relative density of 2.65 is assumed for

Progress has been made on Savage's first point, the question of non-sphericity of particles. It has been addressed by laser scattering techniques for particle suspensions (e.g. Konert & Vandenberghe 1997; McCave et al 1986; Ma et al 2000). This technique has enabled an allowance to be made for particle shape, and has generally led to a small but significant increase in clay fraction estimation. Such an allowance is not specified in SANS 3001 GR2.

Savage's fourth point seems to have drawn little attention, since almost all non-organic soil components have densities reasonably close to 2.7, and the likely error due to this factor is probably quite small. His remaining two points concern dispersion, and obviously merit attention.

many years (e.g. Rodriguez et al 2011; Rolfe and dispersants which have been in use for to soil mechanics continues to use methods soils testing at this stage of development. ticles and may be too expensive for routine on de-flocculation/dispersion using nonand solute system. Experimental research aspects of dispersion for any specific clay and it remains very difficult to assess most itself as the instrument of investigation dispersion continue to use the hydrometer of error likely to be involved in incomplete et al 1960). Attempts to assess the magnitude groups acting as spacers between clay par-2008). Such dispersants produce functional to be concentrated on ceramics (e.g. Al-Lami traditional de-flocculants currently appears well understood (e.g. Robinet et al 2011), ticles suggests that the problem is far from Work on de-flocculation/dispersion relevant theoretical aspects of dispersion of clay par-Research currently being done on the

(Nettleship et al 1997; Rodriguez et al 2011). This paper is primarily concerned with Savage's second point, the dispersion of clay particles. His third point, clay being carried down with larger particles, follows from this as a matter of course.

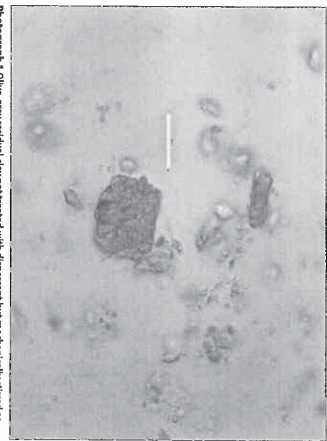
# THEORETICAL BACKGROUND TO DISPERSION OF CLAYS

The behaviour of dispersants is complex and appears to be still imperfectly understood. This outline synthesises information from Das (2008), Zschimmer and Schwartz (2014), Nettleship *et al* (1997) and Robinet *et al* (2011).

particle to peaks on another. adjacent clay particles to maintain mutual electrical attraction by fitting troughs on one ions and troughs between them. This allows surrounding the clay has marked peaks at the on the surface of the clay. The resulting field a clay particle to balance the positive charge tively few of them need to congregate around provide multiple electro-negativity and relatance from the clay surface. Multi-valent ions whose concentration diminishes with discharged ions which form a diffuse layer edges of the clay particles attract negatively ted. Secondly, the positively charged outer increase in volume powerfully when wetapart. This is the reason why some clays can dral sheets of these clays, forcing the sheets etrate between the tetrahedral and octaherior of some types of clay minerals and penattracted by the negative charges in the intecations become mobile. They are strongly surround cations (positively charged metal ions) in the soil. When coated with water the charged, while remaining neutral as a whole), tively charged and the other side negatively (their atomic structure leaves one side posi-Firstly, water molecules, which are polar with water, two things tend to happen. charged. When active clay soils are mixed tend to leave their outer edges positively their inner structure negatively charged and Clay particles carry charges which leave

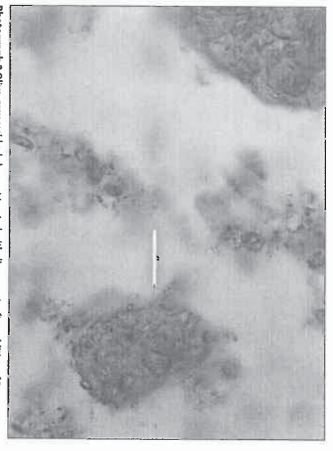
In order to assess clays by their rate of precipitation, as in the pipette and hydrometer methods, it is necessary to disperse the particles of clay into the water through which they precipitate. Mechanical agitation is essential for this, but is not sufficient on its own. Chemical dispersion is needed to break the bonds of electrical attraction holding assemblages of clay particles together.

Dispersants work in three ways. The first is to replace multi-valent ions at the clay surface by mono-valent ions. When an individual clay particle is surrounded by sufficient mono-valent ions to render it electro-neutral, the field surrounding it is relatively uniform; clay particles in such a



Photograph 1 Olive-grey residual clay not treated with dispersant but mechanically stirred as specified in SANS 3001 GR3

Comparison with the 30 micron by 2 micron rectangle suggests that there are particles of about 2 microns attached to several of the silt particles. Some clay-size particles appear to be dispersed in suspension. Note the faint pinkish cloudy patches covering a considerable part of the field of view (most of which is not within the lens's range of sharp focus).



Photograph 2 Olive-grey residual clay, not treated with dispersant, after addition of 3 mg methylene blue per 1 g of soil

Many of the particles of 2 microns and a little larger adhering to the sitt particles show a faint blue outline. Much of the pinkish cloudy area has taken in methylene blue and appears to be composed of extremely small clay particles. Little of the field of view is in focus, but some deeply-stained individual particles smaller than 1 micron are discernible

state cannot attract each other by fitting electrostatic peaks to troughs. The second way is by reacting with multi-valent ions to form chemical complexes, making them unavailable for attraction to clay surfaces. The third manner is by forming functional groups which act as spacers between the clay

particles, effectively preventing them from approaching each other.

The combined action of clay particles, cations and dispersing agents is complex. Above a certain concentration of dispersant the diffused double layer starts to become thinner, repulsion between the particles

reduces, and at higher concentration turns to attraction, allowing flocculation to occur.

Sodium hexamataphosphate is one of the most popular dispersing agents. It is specified in the standards for assessing clay fraction in Britain, America, Canada, Australia, Japan and other countries. It provides mono-valent sodium ions to coat the clay surface, as well as phosphate groups to form complexes with multi-valent cations. Sodium carbonate may be added to increase alkalinity; this has been found to improve the dispersive efficiency slightly in some circumstances (Rofe et al 1960) and to extend the useful life of the dispersant (Nettleship et al 1997).

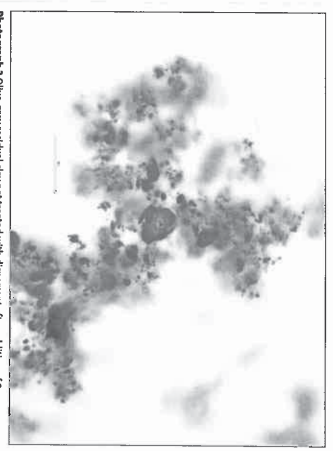
A mixture of sodium hexametaphosphate and sodium carbonate is specified as the dispersant for the hydrometer procedure of SANS 3001 GR3.

## DEACKGROUND AND AIMS

tion indicated by the hydrometer for a range tion projects in South Africa. of clays typical of those found in construcan insight into the reliability of the clay fracsituations. The aim of this study was to gain method may be relevant for a wide range of in South Africa, such shortcomings in the normal for almost all construction projects clay content. Since hydrometer analysis is in the ground surface suggested at least 20% clay on a site where notable shrinkage cracks hydrometer analysis indicated less than 10% indications of clay fraction. In one case technical investigation had given misleading in some of the failures investigated, the geodue to heaving foundations. It appears that bers of failures in government subsidy houses ing solutions to the problem of large num-Investigations are being undertaken by the

adhered to silt and sand particles as conglomerations, while others remained tory. Many clay particles appeared to remain gested that dispersion was not always satisfac of clays using method GR3. The results suginvestigate the efficiency of the dispersion microscope/digital camera combination to to examine the possibility of using an optical the geotechnical research laboratory of CUT series of exploratory tests were conducted at not replicate conditions in the hydrometer. A ing with gold (Nettleship et al 1997). This does treated by techniques such as drying and coatfication. Preparation involves samples being to examine clays with an electron microscope. This has the advantage of very high magni-It is common practice among researchers

The procedures used by soil science laboratories differ somewhat from those for engineering materials. Previous cooperation



**Photograph 3** Olive-grey residual clay not treated with dispersant after addition of 8 mg methylene blue per 1 g of soil

None of the pinkish cloud remains, it appears to have taken in dye and can be deduced to be an aggregation of very small clay particles. Almost all of the silt particles appear to be completely covered with clay. The assembly moves as a flexible unit in the currents caused by evaporation around the edge of the cover slip. No individual clay particles, which are not part of an aggregation, can be seen.



Photograph 4 Olive-grey residual clay treated with dispersant and stirred as specified in SANS 3001 GR3

Many clay-size particles are dispersed but some remain attached to silt particles or form associations of three or more particles not attached to silt.

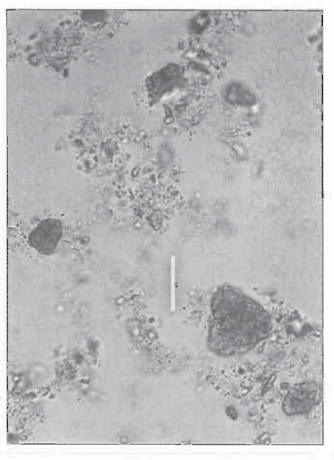
with the Soil Science Department of the University of the Free State had sometimes found higher clay fractions indicated by their procedures. It was arranged for six samples to be prepared by the UFS soil science laboratory using their normal method. The dispersant is 50 g per litre sodium hexametaphosphate solution (the amount

applied depends on soil type), sonification in a dismembrator is specified for clay soils, and mechanical dispersion is of shorter duration but at a higher speed than SANS 3001 GR3. The samples were examined to determine whether dispersion by this treatment was visibly more effective than SANS 3001 GR3 for these six soils.



Photograph 5 Olive-grey residual clay treated with dispersant and methylene blue

The small agglomerations of clay-size particles show little, if any, staining, suggesting very low CEC. The silt particle at bottom centre appears to be coated with very small, high CEC particles which are very darkly stained. The deeply stained agglomeration in the centre is about 50 microns in length and 25 microns in width. It appears to be made of very small, high CEC particles, and it seems to engulf several silt and clay-size particles. Such agglomerations were not uncommon in this sample, but probably not common enough to ensure a meaningless clay fraction from hydrometer analysis.



Photograph 6 Red-brown soil from Limpopo after mechanical stirring without addition of dispersing agent

Many of the sllt-sized particles appear to have clay-sized particles adhering to them. A faint cloudy pinkish haze, as noted in Photograph 1, is again evident. Most of the sllt particles appear to be clustered together in loose associations.

# EQUIPMENT, MATERIALS AND THEIR USAGE IN THE INVESTIGATION

### Microscope and camera

An optical microscope with objectives of 10x, 40x, 60x and 100x was equipped

with a digital camera (resolution 9 mega pixels). Magnification resulting from the combined effects of the microscope's lenses and the camera was assessed by measurements on a 100 lines/mm diffraction grating.

A drop of sample prepared for hydrometer analysis was placed on a microscope slide, covered with a cover-slip and photographed at various magnifications.

Photographs were taken at various locations on the slide. Most of the photographs in this investigation were taken using the microscope's 40x objective since more powerful lenses give a very small depth of focus.

#### Magnification

2 microns respectively. of each rectangle represent 30 microns and indicate the scale. The length and breadth article have a rectangle superimposed to known size. All of the photographs in this is by incorporating a reference object of convenient way of indicating magnification on the photographs produced by the diffraction grating corresponds to 150 pixels Alternatively, the 10-micron spacing on the This implies a magnification of 4 000 times on the screen when using the 40x objective. on the diffraction grating, spaced at 40 mm images showed lines spaced at 10 microns tion can be defined in different ways. The The combined optical and digital magnificacamera using the same objective. The most computer screen that was used to view the

### Variations in procedure

specification is adequate. Examination of the and this part of the investigation served phate (Mishra et al 2011). concentrations of sodium hexametaphos-American standards, which specify different yield for certain clays using the Japanese and by the finding of a difference in hydrometer concentration of dispersant was prompted only to verify whether this aspect of the involved are within these specifications, of soaking and agitation. All of the times Method GR3 specifies only minimum times of dispersant and volume of dispersant used dispersant, time of agitation, concentration examine the influence of time of soaking in ations to the normal procedures in order to Samples were also prepared employing vari-

#### Methylene blue

In addition, samples were treated with methylene blue (MB), with the aim of labelling clay particles for positive identification. Methylene blue (C<sub>16</sub>H<sub>18</sub> N<sub>3</sub>SCl) is an effective indicator of clay, as it readily exchanges places with cations in the clay mineral structure, the amount depending on the cation exchange capacity (CEC) and specific surface area (SSA) of the clay minerals (Turoz & Tosun 2011). Active clays like montmorillonite have high CEC and SSA, and readily take in methylene blue. When MB is available in large concentrations,

considerably more dye. would be coloured only after the addition of low CEC / low SSA clays (e.g. kaolinite) oured after adding very little dye, whereas clays (e.g. montmorillonite) would be colnot be coloured, and high CEC / high SSA could be expected that silt and sand would therefore added to the dispersed samples. It small quantities of methylene blue were the effectiveness of the dispersant. Only the cation balance will inevitably influence additive to the soil solution which affects are, in fact, composed of clay minerals. Any be seen adhering to silt and sand particles whether the clay-size particles which can in a sample, and can also help to establish cation of the types of clay mineral present amounts of dye can therefore give an indideeply stained. Progressive addition of small high CEC/SSA fractions present are already and SSA and show little colouring until clay minerals like kaolinite have low CEC and appear in photographs as dark blobs in which no structure can be seen. Inactive such clays rapidly become totally opaque

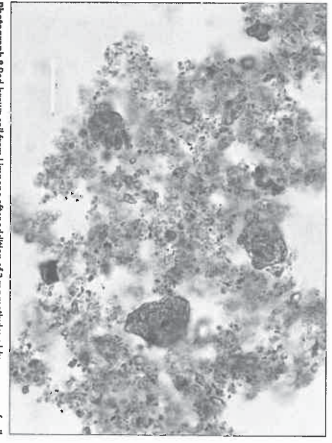
# THEORETICAL CONSIDERATIONS AND STRENGTHS / WEAKNESSES OF THE METHOD EMPLOYED

need dispersion. which do not flocculate and should not question of whether they are clay particles ous range of 1 to 2 microns, raising the particles observed were in this ambiguattempt to establish only particle sizes, not change) depend on clay mineral content, aspects of soil behaviour (e.g. volume mineral classification. Certain important size classification may not correspond with which need to be dispersed, or silt particles mineral content. Many of the individual while hydrometer and pipette analyses Kirsten (1991). There is thus a range where electron micrographs by Bühmann and haloysite, may have particles considerably about 0.1 micron to slightly more than 2 larger than 2 microns, as can be seen in et al 2011). Some clays, e.g. kaolinite and from about 1 micron upwards (Robinet microns; non-clay particles typically range tions of clay minerals typically range from of silt-size. But particles and agglomeraother arbitrary figure of this order) to be from 2 microns to 60 microns (or some smaller to be clay-size particles, and those consider all particles of 2 microns and Soil mechanics and soil science generally

The magnifications possible with the optical microscope and camera combination used in this investigation are probably sufficient to distinguish most of the range typical for clay through silt to sand, but not



**Photograph 7** Red-brown Limpopo soil as in Photograph 6 after stirring without dispersant The pinkish haze and loose groupings of silt particles are again evident. The large silt/sand particle appears to have lost part of its coating of clay-size particles.

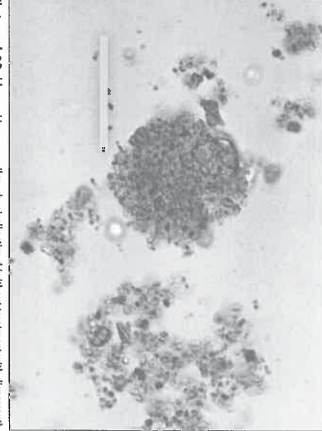


Photograph 8 Red-brown soil from Limpopo after addition of 3 mg methylene blue per gram of soil

The pinkish cloud appears to have absorbed methylene blue, revealing itself to be an extensive agglomeration of small clay particles, as in the case of the olive-grey clay in Photographs 1, 2 and 3. Most of the silt grains which appear to be covered with clay-size particles have scattered spots of dark blue stain suggesting a few high CEC clay particles among many low CEC particles.

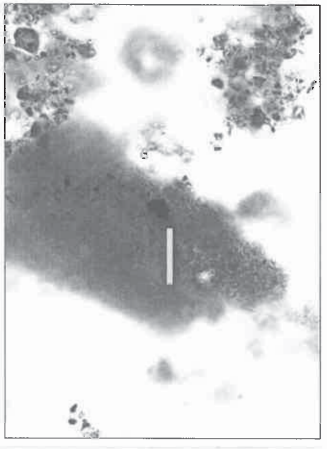
adequate to measure the smallest particles in this range. Since all samples remained in aqueous suspension, all of the smaller individual particles were subject to Brownian motion. At the highest magnification (100x objective – 37.5 pixels per micron, 10 000x magnification on the computer screen), particles at the lower end of the clay-size range could be distinguished in many of the samples, but their Brownian motion

hindered observation or measurement since they suddenly appear in the focal plane, and disappear as they move away from the focal plane. Photographing them was not very successful, possibly because the exposure time of the camera/computer combination was too long. Many small particles were visible and could be photographed where they formed part of large agglomerations or were attached to silt or sand particles.



Photograph 9 Red-brown Limpopo soil mechanically stirred (without treatment with dispersant) and subsequent addition of 6 mg of methylene blue per gram of soil

The dense mass of deeply stained clay particles appears to almost completely engulf a silt particle covered with barely stained clay-size particles.



**Photograph 10** Red-brown Limpopo soil not treated by dispersant with 6 mg/g methylene blue added

The large blue structure is more than 150 microns by 50 microns in size. Within this structure silt particles can be distinguished. Much of the agglomeration appears to consist of clay particles of about 1 micron or smaller. The agglomeration slightly above and left of centre appears to consist almost entirely of a different species of clay particles of about 2 microns which are very lightly stained. Very few soil particles are visible which are not part of an agglomeration.

# **GENERAL CONSIDERATIONS**

The following considerations in terms of the microscopic investigation should be noted:

1. Those samples extracted for microscopic investigation at the UFS laboratory were taken by pipette after a settling time calculated to give only silt- and clay-sized particles at the depth of extraction (larger

particles having settled below this level). The largest particle sizes measured were of the order 50 microns, suggesting that the sample was, indeed, restricted to clay and silt-sized particles. Samples prepared in the CUT laboratory were taken immediately after agitation, and some samples contained particles considerably larger

- than 50 microns, allowing examination of sand grains as well as silt.
- 2. The cover slip over the sample was supported by the largest particles, and consequently a depth of about 50 microns was filled with suspension for the UFS samples, and up to about 100 microns for the CUT samples. Depth of sharp focus at high magnification is far smaller than this and consequently photographs necessarily had most of their field out of focus.
- 3. Since clay sizes range from 2 microns downwards, the concentration of suspension specified in the hydrometer method allows too many clay particles in a depth of 50 microns for convenient optical differentiation. This made dilution of the hydrometer samples necessary. The majority of samples were diluted with three times their own volume of de-ionised water. This dilution was arbitrarily chosen and was considered adequate for this purely qualitative investigation.
- 4 The gap of approximately 50 to 100 the observation of each slide. results in a very limited time available for other clay particles. This consideration particles or formed agglomerations with those which were attached to silt or sand were dispersed and free-floating, and ing between clay-size particles which sible to seal around the edge of the cover by evaporation was helpful in distinguishfound that the movement of water caused slip and prevent evaporation, but it was water around the edges. It may be posallows evaporation of the suspension's microns between slide and cover slip
- 5. When samples dry out they conglomerate, making it difficult to draw conclusions about the behaviour of the clay in conditions relevant to the pipette and hydrometer tests. Only observations in suspension conditions were considered in this investigation.

### INVESTIGATION INVESTIGATION

Samples of six widely different clay soils (from the Free State, Northern Cape, Western Cape and Limpopo) were mechanically agitated, as in procedure GR3, but without first soaking in dispersant. Samples of the same soils were prepared with both dispersant and mechanical agitation at CUT, as per SANS 3001 GR3, and at the Soil Science Laboratory of UFS using standard soil science procedures. From the six soils, two were selected as showing typical features and illustrating the general effectiveness of the investigation's procedures. One soil appeared to show fair dispersion, the other

very inadequate dispersion. The first of these samples is shown in Photographs 1 to 5, the second in Photographs 6 to 13. They provide a reference frame and show widely different clays with and without dispersant, both with and without methylene blue. Features of some of the other soils are shown in the remainder of the photographs.

#### OBSERVATIONS

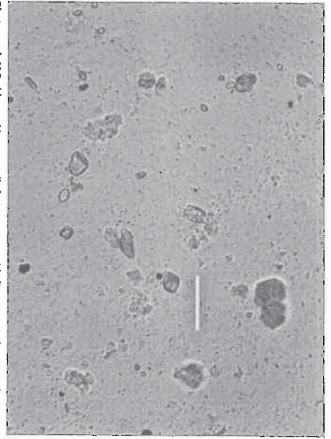
Photograph 1 is of an olive-grey residual clay from a proposed housing development in Bloemfontein. Tests at the UFS Soil Science Laboratory gave LL 72, P1 26 and clay fraction 41% (by both hydrometer and pipette methods). Photograph 2 shows the same sample after the addition of 3 mg of MB per gram of soil. Photograph 3 shows the same sample after the addition of a further 5 mg/g of MB dye.

It appears that a large number of very small clay particles bind considerable numbers of various kinds of particles into associations. Currents caused by evaporation of the suspension's water show that these associations are flexible, but strongly tied together and move as a unit.

Photograph 4 shows the same soil after treatment with dispersant as specified in SANS 3001 GR3. Comparison with Photograph 1 shows a very large increase in clay-size particles dispersed throughout the water. There are, however, some clay-size particles adhering to silt particles, and a number of small agglomerations of clay-size particles with no visible silt core.

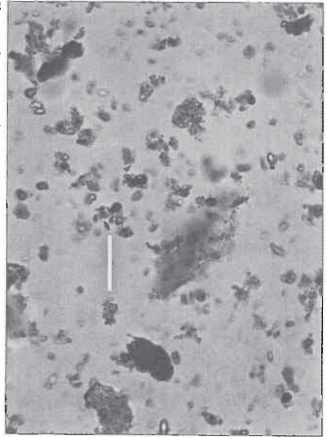
fraction being recorded by the hydrometer. will ensure their contribution to the clay will probably not precipitate at a rate which cles and the agglomeration of very fine clay clay-size particles, the clay-coated silt particlay-size particles. The agglomerations of whether they are free or attached to larger also be seen, and it is difficult to discern stained particles of less than 1 micron can be about 2 microns in size, but small, deeply The majority of dispersed particles appear to it is apparent that dispersion is not complete. dispersion compared to the untreated state, treated sample after the addition of methylene blue. While this sample shows good Photograph 5 shows this same dispersant-

Photograph 6 is of a red-brown soil from the Limpopo Province, which has a history of giving variable results in soil tests and causing difficulties in construction. Treatment was only mechanical stirring of the raw soil without dispersant. Commercial laboratory results for samples sent by CUT as part of a parallel testing programme ranged between 17% and 56% for clay fraction, and between 31 and 43 for PI.



Photograph 11 Red-brown Limpopo soil after treatment with dispersant at the UFS Soil Science Laboratory

Many clay-size particles are well dispersed, but a substantial number remain attached to silt particles. There are also cloudy pinkish areas similar to those in Photograph 1.



Photograph 12 Same sample as Photograph 11 after addition of methylene blue

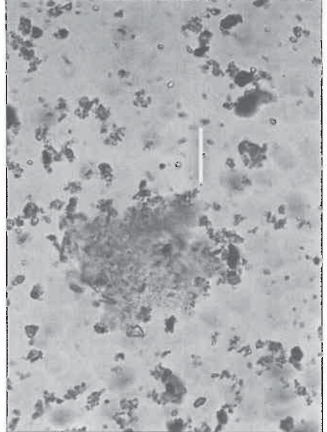
As in the case of the olive-grey Bloemfontein soil there are shadowy pinkish bands associated with the distinctly visible particles. There are also many clay-size particles adhering to most of the silt-size particles.

In Photograph 7 a large grain of silt appears to be mostly covered with clay-size particles. Part of the grain, however, is completely clean and free from clay coating. It is possible that it was struck by one of the paddles of the mechanical stirrer and some of the coating was torn away. The coating of the lower right area seems to have come loose

from the large particle, but remains attached to the clay coating above.

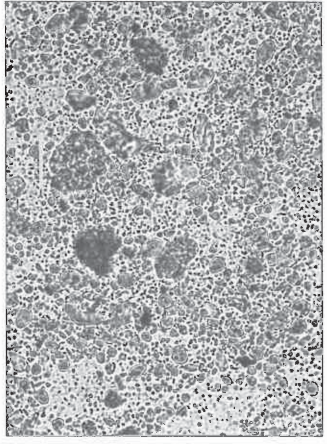
Photograph 8 shows the same sample after addition of 3 mg/g of methylene blue. A few small blue spots are visible on the larger silt-sized particles, but the majority of particles of about 2 microns remain unstained. A clearly visible cloud of very small bluestained particles has largely replaced the faint pink cloud, suggesting that the cloud consists of very small and possibly translucent clay particles with high CEC/SSA.

Photographs 9 and 10 show the same sample after the addition of a further 3 mg/g



Photograph 13 Same sample as Photograph 12

The dispersion and agitation procedures have produced many well-dispersed clay particles, but the large agglomeration shown here is far from dispersed. This sand-size grouping of silt and clay is unlikely to settle at the rate expected of clay, nor are the numerous smaller aggregations.



**Photograph 14** Soil from a subsidy housing project in the Northern Cape after treatment with dispersant and mechanical agitation at the UFS Soil Science Laboratory

The majority of silt particles remain covered with clay-size particles.

of methylene blue. Photograph 9 shows a grain of silt, apparently covered with low CEC clay, almost completely enveloped by high CEC clay particles. A number of small clay particles appear to be dispersed into the surrounding water, but the majority are associated with aggregations. In Photograph 10 a large agglomeration of clay appears to contain several small silt particles of various types.

Photograph 11 shows a sample of the same soil treated with dispersant. Many

small clay particles are dispersed, and, compared to Photographs 6 and 7, the groupings of silt particles have largely disappeared. Yet many of the slit particles remain totally or partially coated with clay-size particles, and many agglomerations of two or more claysize particles can be seen.

Photographs 12 and 13 show the same soil after addition of methylene blue.
Compared to Photographs 9 and 10, where no dispersant was used, dispersion is clearly

improved, but most of the silt particles are seen to be covered with clay, and the faint pinkish clouds again appear to be revealing themselves as very fine clay particles which are not well dispersed and may settle in the hydrometer as silt-size aggregates rather than as individual clay particles.

Photograph 13 shows an exceptionally large agglomeration of small clay particles engulfing several silt particles and numerous 1 to 2 micron clay particles against a background of well-dispersed clay particles. Nettleship *et al* (1997) came to the conclusion that their anomalous observations of settlement in the hydrometer might be explained by agglomeration taking place while particles were settling during the test. It seems more likely that this could be the case here than that such an agglomeration could have survived 15 minutes of stirring at 1 500 rpm after prolonged soaking in dispersant.

The amount of clay which is obviously not dispersed in Photographs 12 and 13 suggests that it is very unlikely that the hydrometer will give a reliable estimate of the true clay fraction of this soil.

Photograph 14 shows a soil from a housing project in the Northern Cape. Hydrometer analysis had shown the clay fraction for almost all of the samples from this site to be very low. This led to a low value of Van der Merwe's predicted heave being accepted for design. Heave damage did, however, occur on the project.

A considerable fraction of the clay-size particles visible in the photograph are attached to silt particles of various sizes. It is not clear whether all the agglomerations of clay-size particles have a silt core, but it appears that much of the clay in this sample has not been dispersed. Hydrometer analysis could underestimate the clay content quite drastically. This might explain the unexpected damage which occurred at the housing project.

Another sample, from less than 100 m away, was prepared according to SANS 3001 GR3 at the CUT laboratory. A house had become structurally unsound due to heave while still under construction a few metres from where this sample was taken. Photographs 15 and 16 are of this sample. In Photograph 15 many of the visible clay-sized particles are attached to silt particles. It appears that the large congregation of claysized particles in Photograph 16 surrounds a root hair or similar thin, thread-like structure

Considerably more clay is undispersed than is dispersed. This would suggest the likelihood of a misleading estimate of clay fraction by the hydrometer method.

Photograph 17 shows a low-activity kaolinitic soil described as "light yellow silty clay" from the Western Cape after

had been incorrectly assessed as silt. the rate expected of silt-size particles and that they had settled in the hydrometer at clay are, however, of silt size and it could be if any, silt is evident. The agglomerations of particles, the clay fraction determination is types. Many agglomerations of clay particles again likely to be unreliable. The description settle at the rate expected of individual clay are evident. Since it is unlikely that these will a range of sizes and probable clay mineral particles; all of the other samples showed tested which showed no small, high CEC clay clay being kaolinite. This is the only sample larger in size, which is consistent with the clay particles are about 2 micron or slightly that its heave potential is very low. All of the It has LL 34, PI 13 and LS 4.6 suggesting dispersion and addition of methylene blue. seems inappropriate, since little,

#### ADDITIONAL TESTS

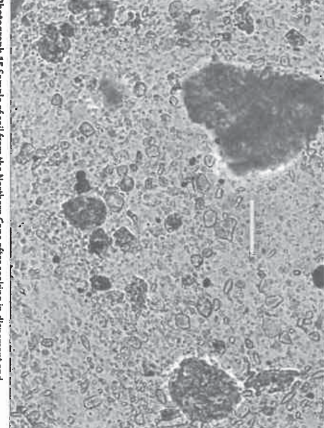
adequacy of this aspect of the specification. and agitation, so this served only to check the includes only minimum times for soaking of mechanical agitation. The GR3 procedure time of submersion in dispersant and time dispersant, volume of dispersant, length of out to assess the effect of concentration of geotechnical research laboratory were carried Photographs 12 to 16). Tests at the CUT Some showed very poor dispersion (as in tions or associations with other particles). no clay-sized material forming agglomerationably satisfactory dispersion (i.e. little or Photographs 4 and 5), none showed unquessamples tested showed fair dispersion (as in Although it appears that one or two of the

SANS 3001 GR3 calls for a minimum of 16 hours submersion in the dispersing agent. Various periods from 16 hours to 2 weeks were tested. No visible improvement in dispersion was observed.

SANS 3001 GR3 calls for a minimum mechanical stirring time of 15 minutes at 1 500 rpm. Various periods from 15 minutes to 24 hours were tried with no visible improvement in dispersion observed. This tends to confirm that the specified minimum times are adequate.

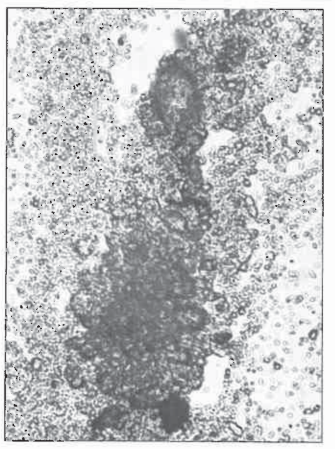
No visible improvement in dispersion was observed by doubling the quantity of dispersant used to treat the samples or by increasing the concentration of dispersant from 40 g/l to 60 g/l. This was not unexpected, since the UFS samples used 50 g/l and showed no visibly significant improvement in dispersion from the GR3 samples.

Samples taken from suspensions permitting little time for settlement, allowed assess ment of the dispersion of clay particles from sand-size particles. Dispersion appeared to



Photograph 15 Sample of soil from the Northern Cape after soaking in dispersant and mechanical agitation as specified by SANS 3001 GR3

This sample was taken a few metres from where a house became structurally unsound due to heave and was demolished before the roof was installed. Soil apparently identical to this from a nearby test pit was assessed by hydrometer as containing only 6% clay. Many apparently clean silt particles can be seen, as can many dispersed particles smaller than 2 microns. The larger particles are all covered with clay-size particles.



Photograph 16 Same sample as Photograph 15 after treatment with dispersant and mechanical stirring

A large number of clay-size particles form an agglomeration around a thin, thread-like structure – possibly a root-hair – while towards the top right many similar size particles appear to be well dispersed.

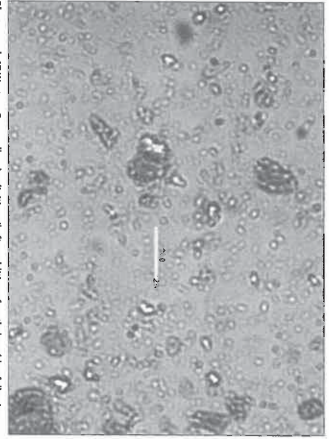
be no better than from silt, as can be seen in Photograph 18.

With particles as large as this, the depth of the suspension between slide and cover slip is so great that very little of the suspended material is in focus. It appears that treatment with dispersant and subsequent

mechanical agitation had failed to dislodge clay particles from the sand grain.

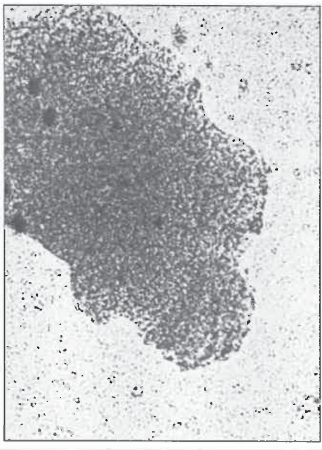
#### DISCUSSION

All of the clays tested showed some lack of dispersion. Every sample showed instances



Photograph 17 Western Cape yellow kaolinitic soil after addition of methylene blue following **SANS 3001 GR3** treatment with dispersant and mechanical stirring to the specification of

Almost all of the particles are about 2 microns or slightly larger. Many form groups of from a few to many particles. Many appear to be properly dispersed, but many are not.



**Photograph 18** Dark-brown sandy clay from a road project at Thaba Nchu in the central Free mechanical agitation according to SANS 3001 GR3 State after preparation for hydrometer analysis by soaking in dispersant and

This sand grain is of the order of 200 microns long and 100 microns wide. It appears to be completely covered with clay-size particles. Many clay particles around it are dispersed, although most are out of focus and cannot be clearly seen.

of clay remaining attached to larger particles or forming agglomerations with other clay particles.

In some cases the lack of dispersion was fairly small, but in some cases a substantial fraction of the clay particles appeared to be undispersed. This suggests that it will not be reasonable to look for some universal factor

by which hydrometer results could be corrected. It appears that predictions based on clay fraction determined by the procedure of SANS 3001 GR3 may be very unreliable for some soils. Since the SANS 3001 procedure is quite similar to that of many other countries, it is likely that this problem may be widespread. The samples prepared at

the UFS Soil Science Laboratory, using soil science procedures with some differences to those of SANS 3001 GR3, showed visibly similar results to those prepared at the CUT laboratory using the GR3 procedure.

results for these soils. several types of clay, it will not give reliable kaolinite). If a dispersant is not efficient for large, low CEC particles (much of it probably (much of it probably montmorillonite) to of clay ranging from small, high CEC particles since most of the soils tested showed mixtures Changing dispersant may be futile, however, soils and completely inadequate for others. appears to be reasonably adequate for some dispersant specified for all South African soils Perhaps it is not surprising that the single soils, but is not effective for dispersing lowium hydroxide is very efficient for high-CEC of soil. Rodriguez et al (2011) found that lithing different de-flocculants for different types different types of clay in hydrometer tests. yield given by a number of dispersants across found considerable difference in the clay CEC electropositive soils. Rolfe et al (1960) It might be reasonable to consider specify.

The methods of quantitatively assessing the efficiency of de-flocculants for geotechnical and soil-science purposes (Rodriguez et al 2011; Rolfe et al 1960) take hydrometer yield as the standard of comparison. There appears to have been no attempt to assess how much of the clay remains undispersed. The use of even the most efficient dispersant for any particular clay may therefore give poor results.

There is also the question of mechanical agitation. Rodriguez et al (2011) noted that horizontal mechanical shaking in helicoidal motion, with the addition of coarse sand as an abrasive, is more effective for dispersion than the conventional method. They did note, however, that its efficiency is not the same for all soils.

#### CONCTANTON O

It appears that Savage's suspicion that "deflocculation of many clays is seldom fully completed at the time of testing" is well founded. None of the clays tested reached good dispersion, even when all aspects of the dispersion procedure were extended substantially. Clay coating of large (silt/sand) particles was observed to some extent in all samples which contained silt and sand particles — in some cases to a very considerable extent. Such particles will probably settle at the rate of silt/sand particles and their clay coating will not be assessed with the clay fraction.

It may be advisable to consider the hydrometer unreliable for any critical

analysis such as heave prediction. Work in progress at the CUT soil mechanics research group is attempting to find more reliable methods of assessing clay fraction, but this is at a very early stage and is therefore unlikely to be able to give reliable quantitative results for at least two years. The quest for better methods of assessing clay content should perhaps become a priority on a wider scale.

### ACKNOWLEDGEMENTS

The authors wish to express their thanks and appreciation to Prof L van Rensburg and Mrs Y Dessels of the Free State University, and Prof SW Jacobsz of the University of Pretoria, for their assistance and encouragement, as well as the National Research Foundation (NRF) for its financial support of the CUT soil mechanics research group's work.

#### REFERENCES

- Al-Lami, H S 2008. Evaluation of some water-soluble dispersants for different types of kaolin powders.

  Al-Tagani, 21: 95–100. Available at: http://www.iasj.net.
- Bühmann, C & Kirsten W F A 1991. The mineralogy of five weathering profiles, developed from Archaean granite in the eastern Transvaal, Republic of South

- Africa. South African Journal of Plant and Soil, 3: 126-152. DOI: 10.1080/02571862.1991.10634823.
- Das, B M 2008. Advanced Soil Mechanics, 3rd ed. New York: Taylor and Francis.
- Konert, M & Vandenberghe, J 1997. Comparison of laser grain size analysis with pipette and sieve analysis: A solution for the underestimation of the clay fraction. Sedimentology, 44(3): 523–535.
- Ma, Z, Merkus, H. G, de Smet, J. G. A. E., Heffels, C. & Scarlett, B. 2000. New developments in particle characterization by laser diffraction: Size and shape Powder Technology, 111(1-2): 66-78.
- McCave, I N, Bryant, R J, Cook, H F & Coughanowr,
  C A 1986. Evaluation of a laser-diffraction-size
  analyser for use with natural sediments. *Journal of*Sedimentary Petrology, 56(4): 561–564.
- Mishra, A K, Ridlo, M O & Higashi, T 2011.

  Comparison of the clay fraction of Ariake clay determined by Japanese and ASTM standards.

  Marine Georesources and Geotechnology, 29(2):
  110–116. DOI: 10.1080/1064119X.2012.525138.
  29:110-116, 2011.
- Nettleship, I, Cisko, L & Vallejo, L E 1997. Aggregation of clay in the hydrometer test. Canadian Geotechnical Journal, 34: 621–626.
- Robinet, J.C., Coelho, D. & Altmann, S. 2011. State of the art on cation diffusion in clay mineral systems. Karlsruhe, Germany: European Commission (Community Research). CatClay Report 15/2011
- (Community Research), CatClay Report 1/5/2011.

  Rodrigues, C, de Oliveira, V A, da Silveira, P M & Santos, G G 2011. Chemical dispersants and pretreatments to determine clay in soils with different

- mineralogy, Journal of the Brazilian Society of Soil Science, 35(5): 1589–1596. Available at: http://www.redalvc.ore.
- Rolfe, B N, Miller, R F & McQueen, I S 1960. Dispersion characteristics of montmorillonite, kaolinite, and illite clays in waters of varying quality, and their control with phosphate dispersants. Washington DC:

  Geological Survey and Colorado State University,
- SANS (South African National Standard) 2011. SANS 3001: 2011, Edition 1.1. Civil Engineering Test Methods, Part GR3. Pretoria: SABS Standards Division.

Professional Paper 334-G

- Savage, P F 2007. Evaluation of possible swelling potential of soil. *Proceedings*, 26th South African Transport Conference (SATC 2007), Pretoria, 9–12 July 2007.
- Skempton, A W 1953. The colloidal "activity" of clays. Proceedings, 3rd International Conference on Soil Mechanics, Zurich, J. pp 57-60.
- Turoz, M & Tosun, H 2011. The use of methylene blue test for predicting swell parameters of natural clay soils. Scientific Research and Essays, 8: 1780–1792.
- Van der Merwe, D H 1964. The prediction of heave from the plasticity index and percentage clay fraction in soils. *The Civil Engineer in South Africa*, June: 103–107.
- Zschimmer & Schwarz GmbH & Co 2014. Mechanisms of action of deflocculates and dispersants in ceramic bodies. Ceramics. Available at: http://www.zschimmer-schwarz.com/en/simon/zschimmer-schwarz/media/site/downloads/fachinfo/eng/Fach/E\_WirkmechVerfluess.pdf.