

RESIDUAL STRESSES IN D/H DRILL BITS

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THE APPLICATION OF
X-RAY DIFFRACTION AND FINITE ELEMENT ANALYSIS
TO THE DETERMINATION OF RESIDUAL STRESSES AND STRAINS
ON THE SURFACES OF BRAZED COMPOSITES OF STEEL
AND SINTERED TUNGSTEN CARBIDE-COBALT COMPOSITES

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by

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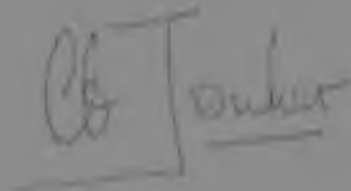
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UNDERTAKING

I hereby certify that this is my own
work and has not been submitted for a
Doctor of Philosophy degree at any
other University

A handwritten signature in cursive script, appearing to read "C. G. Jonker". The signature is enclosed within a rectangular box that has been hand-drawn.C. G. JONKER

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SYNOPSIS

The strains on the surface of the components of a cruciform 'down-the-hole' bit were investigated because of the importance of the surface condition under fatigue loading. X-ray diffraction was selected as the most suitable technique as it may be used to measure elastic strain separated from plastic strain and to determine stress gradients over small distances. Stress values calculated from X-ray line displacement measurements on the (211) d-spacing of hardened IN 100 steel were found to correlate directly with those obtained by strain gauge measurement.

X-ray diffraction peak profile analysis was employed to determine values of the elastic strains, line broadening as a result of plastic strains and the particle size of tungsten carbide grains on the surface of sintered tungsten carbide-cobalt compacts. The surface strains and particle size resulting from several different methods of surface finishing operations were determined. The annealing process implicit in the drill re-boring operation was found to remove the greater part of the surface residual stresses.

Residual stresses in the drilling rate of a cruciform 'down-the-hole' drill bit were investigated using an enclosure, made of the same materials as the drill bit and having a similar geometry. The symmetry of the geometry of the cruciform bit permits the prediction of the residual stress pattern in the entire enclosure virtually by finite element analysis using the enclosure geometry, material constants and thermal history. The residual stress data calculated from X-ray diffraction measurements correspond with the residual stress pattern evaluated by finite element analysis.

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This stress pattern is presented for one sector of the analogue by loci defining the magnitude, type and direction of the principal and shear stresses. This close correspondence between the experimentally determined and finite element analysis residual stress pattern allows the use of the finite element analysis in estimating the residual stresses in the surface of the sintered carbide compacts and those generated prior to and after the martensitic transformation of the EN 30B steel.

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I. Introduction

1.1 Residual Stresses in 'Down-the-Hole' Drill Bits

Residual, internal or locked-in stresses may be defined as the system of stresses which can exist in a body when it is free from external forces. These residual stresses are to be considered as purely elastic stresses. There are two distinct types of these stresses, a macro- and a micro-stress.

The macro-residual stresses, with which this project is concerned, vary continuously through the volume of the body and act over regions which are large compared with atomic dimensions.

Micro-stresses act over dimensions as small as several unit cells, although their effect may extend throughout most of a grain. A major contribution in this field was made by Lammle and has been reviewed by Nabarro⁽¹⁾. He attributes these stresses to the failure of neighbouring grains to fit together, due to anisotropic thermal expansion. This is true for all crystal structures of low symmetry, in contrast with the high symmetrical cubic lattices, which behave isotropically during thermal expansion.

The residual stresses to be measured in this project occur in 'Down-the-Hole' drill bits, commonly abbreviated to D/H bits. The drilling machine is lowered into the drill hole with this drill, so that the drill bit is very compact, and no extensions are required.

The main components of the D/H bit are a steel shaft and crown; in the crown four hard-metal cutting faces are located in cruciform positions (See Figure 1.1).

The basic function of the shank is as a connection with the drilling machine, and it is designed such that the power-drive can rotate the D/H bit in conjunction with the percussive motion. This motion is transmitted to the drill bit via splines machined on the shank, on which the power drive is located.

The crown tapers outward from the shank to the desired gauge width and consists of four wings separated by flutes. Each wing contains one radial slot to accommodate a hard metal insert. The smaller D/H bits have wings at angles of 90° to each other, while the larger units are in the shape of a cross with an acute angle of 80° . The flutes provide a cooling effect and allow the removal of grit from the drill faces. The hard metal inserts are located in the centre of each wing, radiating out from the central ferrule. The wings are tapered from the flutes towards the inserts, so that the latter occupy the extreme positions in the drill face and can act as cutting faces.

The steel body of the D/H bit is turned, drilled, milled and machined to the right dimensions prior to the brazing of the hard metal inserts into the wings. Depending upon the type of D/H bit, flushing channels are drilled into the body. These flushing channels consist of one central channel through the shank into the crown, where four other channels radiate out into the four flutes or the wing faces, providing internal cooling of the drill by air or water.

The complexity of construction of the crown, i.e., the presence of flutes and flushing channels in the steel, in addition to the hard metal inserts, which are brazed into the wings, indicates the existence of many variables which may influence the service life of a D/H bit.

The loading condition of a D/H bit during its service life consists of fatigue loading perpendicular to the face of the drill bit and a radial compressive stress acting on the sides of

the crown of the drill bit.

Fatigue failure is usually initiated at the surface of an object under load and hence the surface condition of the bit is of extreme importance. Notches present by virtue of the geometry of design, surface roughness and surface residual stresses may affect the fatigue life of the drill bit considerably. Surface roughness is frequently an adverse factor in a sample exposed to loading, as it introduces numerous physical defects which seriously reduce the fatigue strength by acting as stress raisers. Surface residual stresses, as introduced by shotpeening, grinding and quenching after heat-treatment can either contribute to or detract from the service life of a drill bit. Shotpeening usually introduces a favourable compressive residual stress pattern, but excessive shotpeening damages the surface, introducing localized stress raisers and reducing the fatigue performance. Tarasov, Tyler and Leiner⁽²⁾ have found that grinding of hardened steel can produce either tensile or compressive surface residual stresses, depending upon grinding conditions.

In most fatigue investigations, it has been recognized that compressive residual stresses in the surface are favourable factors for an increased fatigue life, hence it is very important to assess whether compression or tensile residual stresses are present in the surface of a drill bit. The final grinding of the cutting faces in the crown of a D/B bit after the brazing operation will usually introduce compressive residual stresses. However it is possible to initiate subsurface fatigue failure if a very steep stress gradient exists leading towards a tensile residual stress below the surface. It is therefore extremely important to determine the residual stress pattern just below the drill bit surface underneath the stress pattern introduced by grinding. Any residual stress present not originating from these machining stresses is due to thermal shrinkage effects between the hard metal inserts, brazed joints, and steel sections of the crown. These residual stresses are generated by

differential thermal contraction of the components of the drill crown when cooled to room temperature from the brazing temperature.

During manufacture of cemented carbides from their chemical constituents in powder form, several restraints in the body of the compact generate residual stresses. These are the following:

1. Thermal stresses right throughout the body of the compact caused by the differential shrinkage of the cobalt binder and the tungsten carbide grains.
2. Mechanical stresses in the surface layers of the compact, caused by grinding and shot-blasting operation during manufacture. These latter stresses are partially relieved by heat-treatment during D/H Bit assembly, but are of great importance in relation to the brazing characteristics of the hard metal compact.

After brazing of the inserts into the slots of the drill bit crown, further grinding stresses are introduced in the surface of the hard metal compacts by the final sharpening operation. These stresses will be present in the cutting surface only, and will modify any residual thermal stresses.

During the loading cycle of the brazing operation the following series of events generate residual stresses near the tungsten carbide-binder metal and steel/braze metal interfaces.

- a) The tungsten carbide inserts are brazed into the steel slots at a temperature of 870°C . The braze metal solidifies over the temperature range $650^{\circ} - 634^{\circ}\text{C}$. When at 634°C all three different components of the group are molten, interaction is expected to occur between them upon cooling to room temperature, due to their different specific thermal contractions. The macrostresses so generated in the braze, the sintered carbide and the steel are only partially relieved by

plastic / - 5 -

b) plastic deformation of the relatively soft braze alloy. The steel section of a D/H bit is made from EN 30B steel, which is an airhardening steel. After austenitization of the steel at 870°C , the brazing temperature, the steel is martempered, this heat-treatment being described in Section 3.1. The austenitic structure transforms to martensite and possibly some lower bainite, this transformation generating volume-stresses inside the steel, which are part of the residual stress pattern in the composite D/H drill.

A survey of the modes of failure of all drills returned over a period of four months indicates that more than sixty per cent of all failures were initiated at the braze-steel and braze-sintered carbide interfaces and had propagated through either the steel wing or the sintered carbide insert. A smaller number of drills had failed by excessive shank and spline wear and failure of the braze metal insert.

It was therefore decided to select and establish a stress measurement technique with the intention of determining the magnitude, type and distribution of the residual stresses in the surface layers of the drillhead, with the ultimate aim of investigating the possible connection between residual stress and failure of the drill.

Major residual stresses in the surface of a D/H bit with a right angle crossform geometry are generated during cooling from the brazing temperature, and are increased in addition by grinding and machining operations on the surface of the tungsten carbide-sintered hard metal inserts and the steel crown of the drill bit respectively.

1.2 Selection of Method of Stress Measurement

All stress measurement methods are based upon experimental measurement of the corresponding strains.

Several methods of measuring macro-residual stresses exist and these fall into three broad classes, i.e., mechanical-, deflection- and X-ray techniques.

Using mechanical methods the residual stresses are calculated from the measurements of strain that are obtained when the body is sectioned and the locked-in residual stresses are progressively released. These strains may either be measured by electrical resistance strain gauges or by a micrometer. This technique is rather laborious and therefore methods which are more rapid have been developed. These methods involve the mechanical slitting of the specimen and measurement of the deflection of the slit element.

The X-ray method was considered the most suitable for the present project and was selected for the following reasons:

- a) The X-ray method is non-destructive. The extreme hardness of the sintered tungsten carbide compacts, and to a lesser degree of the martensitic steel, make mechanical and deflection methods extremely difficult to use. Furthermore, the values of strain measurements by either of these two methods will be the average of strains over the whole sintered tungsten carbide specimen. This renders these methods useless, since the components of a P/B have reacted to different extents to these techniques of stress relief.
- b) The X-ray method can determine the residual stress over very small regions of approximately 5 square millimetre in area by reducing the cross-section of the X-ray beam to a circular beam approximately two millimetre in diameter.

This feature makes the technique particularly suitable for measurements of steep stress gradients extending over a limited region.

- c) The X-ray method makes it possible to measure elastic strains directly and to measure line broadening of diffraction peaks yielding information on plastic straining.

1.3 Development of the X-ray Strain Measurement Technique

The underlying principle of the method is the determination of the change in interatomic spacing (d) of a particular set of crystallographic planes when the crystal or polycrystallin aggregate is strained in a homogeneous manner. These atomic strains are calculated from the diffraction data relating to the specific set of crystallographic planes. These diffraction data are obtained from X-ray diffraction peaks, which are a plot of diffracted X-ray intensities versus diffraction angle θ . Monochromatic X-radiation will yield a unique diffraction peak for each set of crystallographic planes and when the crystal is strained the angular shift of such a peak is a measure of the change in interatomic spacing (d).

The initial experimental work establishing this technique was performed using X-ray powder cameras, and the strains were measured from the displacement of the diffraction lines on the X-ray films with respect to the X-ray lines of unstressed material. This technique is only acceptable when the diffraction lines are sharp and well defined, which is not the case for hardened steels.

The development of a diffractometer technique by Christodoulou and Rowland ⁽¹⁾ has promoted the X-ray technique of strain measurement on hardened steels to a practical tool. This method makes use of Geiger and other radiation counters to measure the intensity profiles of the diffraction peaks.

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The relative merits of each technique have been reviewed previously (3,4). The X-ray sub-committee of the Society of Automotive Engineers has also reviewed the diffractometer technique and has consolidated the fundamentals of X-ray strain measurements on hardened steels (5).

Initially most practical use of Christenson's and Rowland's contribution was made by the aircraft industry for the measurement of surface stresses in aluminium alloys and high tensile strength steel components (6,7). Subsequently this technique was used on other steel and pure iron specimens (8,9).

Further sophistication, but no essential improvements, have been developed concerning computerisation of the X-ray diffraction data-collection and evaluation procedure (10).

One further step in the development of this technique was the measurement of stresses within a body, in contrast with the surface stresses considered up till now, and this has been described by several authors (3,11,12). It necessitates the sectioning of the specimens but corrections have to be made for the stress relief caused by removal of layers of material (5).

Some experimental results of internal residual stresses corrected for layer removal stress relief, have been published (11,12). In particular the measurement of stresses in carbonised metal surfaces (11).

2. X-ray Diffraction Technique - Theoretical Aspects

2.1 Principles of X-ray Diffraction Strain Measurement

The diffraction of X-rays from the atomic planes in a crystalline material is expressed by Bragg's Law

$$n \lambda = 2 d \sin \theta \quad (1)$$

where λ = wavelength of incident X-rays
 n = order of reflection, which is always an integer
 d = interplanar spacing of (hkl) planes which are diffracting
 θ = Bragg angle of reflection

This derivation of this equation is described in all standard science textbooks.

Fundamentally the X-ray diffraction technique measures the elastic strain from the change in the average interplanar spacing (d) of a particular set of crystallographic planes (hkl). The plastic strain present is a function of the change in diffraction peak profile when stress is applied. The incident X-ray beam originates as the characteristic radiation of a pure chemical element excited by a high voltage, hence the wavelength, λ , lies in a narrow range. After diffracting from the specimen the beam is further monochromated by a metal filter with a suitable absorption edge wavelength, which filters out the K_{β} radiation plus most of the background radiation. The diffracted X-rays which simply ionize the gas molecules in the counter consist of K_{α} radiation, plus a background radiation of large wavelength range.

The wavelength K_{α} of the element is the value for λ used in Bragg's law, i.e., λ is a constant, which simplifies this law to

$$d = \frac{k}{\sin \theta}$$

where k = constant.

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This relationship indicated that for a small variation in interplane spacing (d) the diffraction angle θ will show the greatest variation for the higher order of (hkl) reflections, i.e., those which have large angles of diffraction. Christenson and Rowland ⁽³⁾ have shown that the X-ray diffraction technique is useful only when diffraction angles in the range $70^\circ \leq \theta \leq 80^\circ$ are used. The maximum value, $\theta = 80^\circ$, which may be used is imposed by the mechanical limitations of the X-ray goniometer. The minimum value of $\theta = 70^\circ$ is chosen so that small changes in (d) may still be associated with appreciable change in diffraction angle θ .

Therefore a set of crystallographic planes and wave-length of radiation are chosen in such a way that the diffraction angles fall within the accepted range, on condition that the peak to background intensity ratio of the diffracted radiation is sufficiently high, i.e., of the order 2 : 1.

The X-ray method, for determining elastic strains only, is fundamentally a measurement of a set of interatomic spacings, which are altered by elastic stresses. The theory of elasticity assumes the material under state of stress is being measured to be elastic, homogeneous and isotropic, assumptions which are usually true in the case of polycrystalline metals.

From the theory of elasticity the following X-ray stress-strain relationship can be derived ⁽⁵⁾,

$$\sigma_\psi = \frac{E}{1 + \nu} \left(\frac{\Delta d}{d} + \frac{1}{\sin^2 \psi} \right) \quad (25)$$

alternatively

$$\sigma_\psi = \frac{E}{1 + \nu} \left(\frac{\Delta d}{d} + \frac{1}{\sin^2 \psi} \right) \quad (26)$$

and the corresponding strain

$$\epsilon_\psi = \frac{\Delta d}{d} + \frac{1}{\sin^2 \psi} \quad (27)$$

where $\psi = 45^\circ$

Where d_ψ and d_ϕ are the interatomic spacings of the planes (hkl), which respectively subtend angles $\psi \neq 0^\circ$ and $\psi = 0^\circ$ to the specimen surface and σ_ϕ and ϵ_ϕ are respectively the stress and strain in direction ϕ in the surface of the specimen.

Often, when X-ray diffractometers are used instead of cameras, it is convenient to express the stress in equation (2b) in terms of angle 2θ rather than interplanar spacing d .

To this end, differentiating Bragg's Law (equation 1) yields

$$-\frac{\Delta d}{d} = -\cot \theta \cdot \Delta \theta \quad (4)$$

which is equivalent to

$$\frac{\Delta d}{d} = -\cot \theta \cdot \frac{\Delta 2\theta}{2} \quad (5)$$

Now $\frac{\Delta d}{d} = \frac{d_\psi - d_1}{d_1}$ and $\Delta 2\theta = 2\theta_\psi - 2\theta_1$

and substituting these relationships in equation (3) yields

$$\sigma_\phi = \frac{(2\theta_1 - 2\theta_\psi) \cdot \cot \theta}{2} \cdot \frac{E}{1+\nu} \cdot \frac{1}{\sin^2 \psi} \quad (6)$$

If the stress is measured on a hardened steel, and the same angle ψ is used in every measurement, then all the terms on the right hand side of equation (6) except those involving the θ angle, are constant, so that equation (6) may be written as

$$\sigma_\phi = K \cot \theta (2\theta_1 - 2\theta_\psi) \quad (7)$$

where θ is expressed in radians.

Because the variation in θ for any one material is very small, $\cot \theta$ may also be considered a constant, hence

$$\sigma_{\phi} = K (\epsilon_{\phi} - \epsilon_{\psi}) \quad (8)$$

the constant K being the stress factor. It is convenient to determine this constant K during calibration experiments, so that the mechanical elastic constants E and ν do not have to be used when calculating stresses from X-ray strain measurements.

2.2 Mechanical Elastic Constants versus Stress Factor K

Any conventional (other than X-ray method) strain measurement method will yield values of strain averaged over all the grains and all the crystallographic planes in the direction of strain measurement. In contrast, the X-ray technique measures the strain on a unique set of crystallographic planes in grains with a specific orientation.

Therefore calculations of stress in a body from X-ray diffraction data using the stress factor K are only comparable with values of stress from conventional techniques using mechanical elastic constants like Young's Modulus, E , and Poisson's Ratio, ν , if the material under examination is completely isotropic. This is not the case for most crystalline materials. Iron for instance is anisotropic and its elastic properties vary with crystallographic direction.

Account may be taken of this anisotropic effect by calculating separate stress factors for specific crystallographic planes from a correlation of the interplanar spacing strain and the average strain as measured respectively by the X-ray method and a conventional technique using mechanical elastic constants.

2.3 The Angular Factor of Diffraction Data

Bragg's law (equation 1) predicts a unique diffraction angle for each set of crystallographic planes, when using monochromatic radiation. The deviation from this uniqueness as

observed for diffraction traces of a polycrystalline aggregate is due to the following factors:

- a) The range of orientation of the grains in a polycrystalline aggregate causes a variable reaction to an applied stress from grain to grain, resulting in a range of diffraction angles.
Consider the unique set of crystallographic planes (hkl). A uniaxial stress imposed on the aggregate will cause strains on the (hkl) planes, which vary in a limited range from grain to grain, depending upon their orientation in the aggregate. A single crystal, elastically stressed in the annealed state, will yield a (hkl) diffraction line for a unique Bragg diffraction angle θ . The polycrystalline aggregate also elastically stressed in the annealed state, will diffract the X-ray beam over a small range of angles. This can be represented by a distribution curve of reflected intensity versus diffraction angle θ , commonly taken as either a Gaussian or Cauchy distribution curve. The specific diffraction angle associated with such a distribution and taken as representative of the (hkl) planes is the diffraction angle θ_p of the apex of the peak which approximates the maximum intensity section of the distribution curve.
- b) Very small crystallites have insufficient resolving power and tend to broaden a diffraction line. The extra situation arises when the crystallites are so fine that the distance between the diffraction planes is comparable with the wavelength of the incident beam and will not give any defined diffraction lines. The broadening due to the particle size effect is completely symmetrical about the position of the line which would be obtained if the particle size were infinite.
- c) The X-ray source used is never perfectly monochromatic, i.e. the small energy spread of the incident X-rays produces a small range of diffraction angles.

$$\theta = \arcsin \frac{\lambda}{2d} = \arcsin \frac{\lambda}{2a \sqrt{h^2 + k^2 + l^2}}$$

- d) When a stress is imposed on a polycrystalline aggregate plastic deformation is present, associated with the elastic strain if the proportional limit is exceeded. This may be due to local stress concentrations, which, when an external stress is applied, can result in localised high stress levels resulting in plastic deformation. Plastic strain causes broadening of the base of the original sharp diffraction peak, and reduces its peak to background intensity ratio, while an elastic strain shifts the diffraction peak profile by interval $\Delta\theta$.
- e) The austenite to martensite transformation is the main contributor to the line broadening phenomenon in the hardened steel. This can be ascribed to the microscopic residual stresses as well as the high dislocation content associated with the martensite transformation. This low temperature transformation reaction is a diffusionless process. The face centred cubic austenitic crystal structure transforms to a body centered structure, but since the amount of carbon present is one hundred times more than can be held in solid solution, this carbon super-saturated body centered cubic lattice is distorted to a tetragonal structure, characterised by severe lattice strain.
- f) Instrumental broadening. The response of the proportional counter and other electronic parts introduces deviation from unique diffraction angles.

3. X-ray Diffraction Technique - Practical Aspects

3.1 Location and Construction of Diffraction Peaks

The accuracy of the method depends upon the location and construction of diffraction peaks which is the fundamental step in the stress determination procedure. The work involved in subsequent steps is time-consuming, but basically mathematical.

Christenson and Rowland (3) and others (4,5,9,10) have demonstrated the superior accuracy of a diffractometer for the examination of diffuse diffraction lines. As the present work is concerned with the examination of diffuse diffraction peaks of hardened steel and tungsten carbide-cobalt compounds, the diffractometer was selected as the X-ray tool.

A diffractometer provides the facility for automatic construction of diffraction peaks on a recording chart during a continuous scan over a range of diffraction angles. This method is useful for the determination of the approximate location of a particular diffraction peak. The precision of the method, however, is low due to the time lag introduced by the components intermediate to and including the counter and the recording pen.

The more precise construction method used, is the following:

- a) The specimen is irradiated and diffraction data are collected over the range of diffraction angles $\theta_1 \rightarrow \theta_2$ in which the apex diffraction angle occurs as has been located by the graphical method.
- b) The specimen is irradiated at angle θ_1 for a fixed period of time and the diffracted X-ray intensity is recorded as the number of pulses per unit time.
- c) The diffraction angle is increased by a small increment $\Delta\theta$ and the procedure is repeated until intensity data for all angles in the range are obtained.

By using extended periods of counting time accurate values of the intensities of the diffracted beam can be obtained for each diffraction angle in the range $\theta_1 \rightarrow \theta_2$. Statistically this reduces the percentage error, as it is related to the number of pulses counted by the electronic circuit, and is inversely proportional to the square root of the total number of counts.

The K_{α} radiation of the source has the two distinct wavelength components, K_{α_1} and K_{α_2} , which for a well resolved diffraction trace will occur as two distinct peaks close to one another. The stresses present in the specimen, and particularly the line broadening due to the martensite transformation in steel, cause the K_{α_1} and K_{α_2} peaks to merge in such a way that they cannot be resolved, except by algebraic means. Therefore the merged diffraction peaks are treated as one single diffraction peak of K_{α} radiation and for calculations the K_{α} wavelength, which is a composite value incorporating the K_{α_1} and K_{α_2} wavelengths, is used. The apex angle of this diffraction peak can be calculated algebraically by fitting to a hypothetical parabola the diffraction data for all intensities more than eighty per cent of the maximum intensity recorded (14,15). (See Figure 7). Alternatively the apex angle may be constructed graphically by extrapolating the linear sections on each side of the peak and identifying the angle of intersection as the angle at the apex (3).

The latter method is very useful for sharp diffraction traces, as these exhibit peaks with linear sections of considerable length. The former parabolic method is selected for hardened steel diffraction traces, as these exhibit no distinct linear sections.

For example (Figure 2) for the intensity data for the three peaks θ_1 , θ_2 and θ_3 separated by equal intervals $\Delta\theta$ where θ_2 exhibits the highest intensity value and θ_1 and θ_3 have intensities which are at least 80 per cent of that of θ_2 . Using these data the apex angle $2\theta_p$ is calculated from the following mathematical relation (3), assuming a parabolic diffraction profile.

$$2\theta_p = 2\theta_1 + \sum_{n=1}^N \left(\frac{I_n - I_1}{I_2 - I_1} \right) \Delta\theta \quad (9)$$

where $N = 17$

where $a =$ intensity difference between $2\theta_1$ and $2\theta_2$
 $b =$ intensity difference between $2\theta_3$ and $2\theta_2$
 $c = \Delta 2\theta$

The intensity values used are the periods of time required to accumulate the same number of counts for each angle 2θ .

At this stage it is important to point out that it is not essential to determine the true and absolute angular position of the apex. An arbitrary position may be used, as the strain is only calculated as the difference between two relative apex angles. Hence accuracy of the method will hinge on the precision of repeating apex-positions and not on their absolute positions.

3.2 Correction of Diffraction Data for Intensity Factors Dependent upon θ

A certain amount of asymmetry is introduced into every diffraction line, because the θ dependent intensity factors are closely associated with the diffraction and scattering process.

These factors vary slightly with diffraction angle θ , but become significant in magnitude for diffuse peaks, as these extend over a considerable $\pm \theta$ (diffraction angle) range. All intensity data must be corrected for these factors a priori to obtain a good fit and the location of the apparent peak positions.

These factors are more discussed in detail in standard texts (16,17) and are identified as the Lorentz polarization factor and the absorption factor. The first factor is a combination of several θ dependent factors, while the second is dependent upon the mean path length of the X-rays within the sample and is only a function of diffraction angle θ for all angles ψ about the sample.

Values for these two factors are readily calculated and a combination of these two as a single correction factor was used in this project, as computed by Ricklefs and Shipley⁽⁵⁾. The mathematical expressions for these factors are as follows:

$$\text{L.P.} = \text{Lorentz Polarisation Factor} = \frac{1}{8} \cdot \frac{\sec^2 \theta}{\sin^2 \theta} \quad (10)$$

$$\text{A} = \text{Absorption Factor Relation} = 1 - \tan \psi \cot \theta$$

$$\text{Corrected factor from Ricklefs and Shipley} = \frac{1}{4} (\text{L.P.})(\text{A}). \quad (11)$$

3.3 Focussing Implications of Varying the Angle ψ

For the normal diffractometer geometry, i.e. when $\psi=0^\circ$, the X-ray beam diffracted from the (hkl) planes, will always focus on the circumference of a circle, defined by the goniometer axis as centre, and the source to specimen-surface distance as radius (Figure 3a).

As soon as the specimen surface plane is rotated about the goniometer axis independently of the receiving slit position, i.e. where $\psi \neq 0^\circ$, the rays diffract from the (hkl) planes which subtend the angle ψ to the specimen surface, and will focus at a distance either greater or smaller than the radius of the circle, for respectively clockwise and anticlockwise rotation of the specimen surface (Figure 3b). Hence the standard condition for focusing is not longer at the detector position and consequently the detector distance has to be altered to a different position Δ for every angle ψ .

The proper focussing distance of the receiving slit may be calculated from the following relation:

$$\Delta = D \cdot \frac{\cos(\psi + \eta)}{\cos(\psi - \eta)} \quad (12)$$

where $\eta = \theta - \psi = 0$

D = source to specimen distance

Δ = required focussing distance

4. APPENDIX

4.1 Philips Diffractometer

A standard Philips diffractometer with a long shield vertical anode tube and a vertical goniometer with a horizontal specimen rotation axis was used. The specimen rotation and the counter rotation were geared to a 1 to 2 ratio, i.e. the θ to 2θ relationship.

The counting tube used was a proportional counter, which has the advantage of a linear response between the ionization in the gas chamber and the intensity of radiation for a considerable range of X-ray intensities. The voltage signals from the counter were fed through a voltage discrimination unit before reaching the electronic counting unit.

For steels, a chromium X-ray tube with Vanadium foil filter was selected, instead of the usual cobalt tube iron filter combination, because of the former's superior peak to background intensity ratio of the diffraction peaks of the specific crystallographic planes under examination. The wavelength range of the cobalt tube causes fluorescence of iron K radiation by the specimen, which increases the background radiation level.

For tungsten carbide diffraction, the cobalt tube was selected on the basis of obtaining a diffraction range in the required range $10^\circ \leq 2\theta \leq 160^\circ$ having a satisfactory peak to background intensity ratio.

4.2 Alterations to the Standard Diffractometer

- i) The divergent slit assembly was replaced by a pinpoint collimator of the usual Debye-Scherrer powder camera design. The goller baffles in the divergent slit assembly, which are present for collimating purposes, were also removed.

The collimator γ -20

The collimator reduced the beam cross section from a rectangle of approximately one hundred square millimetres to a circle with a diameter of approximately two and a half millimetres.

The collimator was used primarily to reduce the area from which diffraction data were obtained, and hence to improve the localisation of the strains measured, i.e., diffraction data could be obtained from areas right next to the braze metal, which were not just an average of the strain over the whole adjoining section. The receiving slit assembly was replaced by a unit from which the Soller baffles were removed. This improved the intensity of the diffracted radiation which reached the counting tube. The pinpoint collimator reduced the intensity of the incident beam proportional to its reduction in cross-sectional area to a narrow circular beam. The Soller baffles in the receiving slit assembly were superfluous, as they were intended to collimate the rectangular diffracted beam.

Incident intensity and collimation of the beam were improved by changing the X-ray tube from line to point focus. This was achieved by rotating the central glass portion of the tube, containing the anode, through ninety degrees with respect to its heating. This meant that the X-ray beam from the source was identical with that of the standard Debye-Scherrer powder camera set-up, i.e. radiation from the X-ray tube emerged through the pinpoint collimator now centred from the edge of the anode. This improved collimation considerably as the Debye-Scherrer collimator was designed for pinpoint radiation.

4.3 Attachments - Miscellaneous X-ray and X-ray attachment for the revised Phillips instrument

These two supplementary attachments, manufactured by the Advanced Metals Research Corporation for Norolite, were

required / - 21 -

required to overcome the specimen-size problem and to alter the goniometer for stress determination work.

This metallurgical kit permitted the examination of relatively large specimens, by providing a spacious platform on which the specimen was mounted. This allowed a traverse to be made across the specimen in all directions. To cope with the platform, the goniometer was shifted by 50.8 mm away from the vertical plane of the incident and diffracted X-ray beams.

The stress attachment kit provided a dovetail slide for the proportional counter and receiving slit system, so that the optimum focussing conditions of the diffracted beam could be attained for most angles of rotation ψ .

A four point bending stress attachment was also provided by which a specimen could be stressed in tension or compression while in position for X-ray diffraction. It was used to calibrate the strain as measured by X-ray diffraction against resistance strain gauge measurements.

This bending attachment was used (Chapter 5) to evaluate by X-ray diffraction the elastic constants and the X-ray stress constants of BS 900 steel. These constants were then compared with the elastic constants which were determined by conventional techniques on a macroscopic scale.

The goniometer axis assembly was also altered, to permit rotation of the specimen surface, through the range $-60^\circ \leq \psi \leq +60^\circ$, independent of the counter tube and receiving slit rotation.

4.4 Alterations to the Metallurgical Kit and Stress Attachment

The present project required the handling of large specimens, often 100 mm in diameter as well as the accurate positioning of the specimen with respect to the incident radiation and the provision of the means to collect diffraction

data from identified positions in close proximity to one another. To achieve these aims a lightweight aluminium vice clamp with a maximum gap of 120 mm was used.

This vice clamp was constructed so that it could move along an aluminium dovetail slide and be locked in position. A vernier scale was attached to the vice and the slide to measure the vice movement and hence pinpoint the traverse of the specimen area under irradiation to 0.035 mm when the specimen is clamped in the vice.

The whole unit (see photograph) consisting of vice clamp, dovetail slide and vernier scale was bolted in position on the platform provided with the metallurgical kit.

5. Calibration Experiments on EN 30B Steel

5.1 Specimen Preparation

The surface condition of any specimen to be examined by X-ray diffraction is of paramount importance, because the incident X-ray beam will diffract only from the outermost surface layers. The depth of penetration of the X-rays may be calculated from the following relation (16)

$$I/I_0 = e^{-\mu x} = (K/\rho) \cdot \rho x \quad (13)$$

where μ/ρ = mass absorption coefficient of the metal under examination

ρ = density of the material

x = depth of penetration

I/I_0 = ratio of intensity I at depth x and intensity I_0 of X-rays which strike the surface

For iron, an irradiated material, and chromium $K\alpha$ as incident radiation, the following values are applicable (20)

$$\mu/\rho = 114.0 \text{ cm}^2 \text{ g}^{-1} \text{ and } \rho = 7.8 \text{ g cm}^{-3}$$

Equation (13) / = 23 =

The other diffraction traces, curves b, c, d and e in Figure 4, represent the same {110} diffraction peak after sectioning, grinding and polishing. Ultimately, using an electropolishing technique, the line broadening introduced by the sectioning and grinding operation was removed almost completely, as represented by curve f in Figure 4.

Therefore the outlined specimen preparation technique shows that the initially deformed surface layers may be removed successfully.

The removal of the elastic strains by use of metallographic and electro-etching techniques can be observed only from a comparison of the interatomic or atomic plane spacings which align at the angles $\psi = 0^\circ$ and $\psi = 90^\circ$ to the specimen surface, as discussed in Section 2.2.

These interatomic spacings were measured on several strips of IN 300 steel, which were heat-treated in a manner analogous to that of the actual final heat cycle of a down-the-hole drill bit.

- The heat-treatment of a D/H bit is as follows:
- 1) Annealing at 870°C for 20-30 minutes, followed by cooling to room temperature.
 - 2) Heating to 1000°C, holding for 1 hour, and cooling to room temperature.
 - 3) Heating to 1000°C, holding for 1 hour, and cooling to room temperature.

The final microstructure of the steel is mainly ferritic, but some bainite may be present. The

hardness level is in the range of 44 - 48 Rockwell C.

After heat-treatment, the surfaces of the hardened EN 300 steel strips were prepared as follows:

1. Wet emery paper grind, successively 220, 400 and 600 mesh particle size.
2. Diamond polish ($\phi = 8$ micron) + 10 per cent nitric etch
3. Diamond polish ($\phi = 1$ micron) + 10 per cent nitric etch
4. Electropolish, removing 0.075 mm from the surface.

Table I. presents the data obtained from the four strips by X-ray diffraction. Strips 1 and 2 were prepared up to and including the second step, as outlined above. Strip 3 was prepared up to and including step 3. Strip 4 was prepared up to and including step 4. All strains were measured in the direction of the applied stress, along the longitudinal axis of the EN 300 specimen. The stress and strain values quoted in Table I. were calculated from the double diffraction angle 2θ using equations (1) and (2), viz:

$$n\lambda = 2d \sin \theta \quad (1)$$

where λ = wave length of chromium $K_{\alpha} = 229.1$ nm.

$$\text{hence } d = \frac{229.1}{2 \sin \theta} \quad (2)$$

$$\text{and } \sigma_{\phi} = \frac{d_{\phi} - d_0}{d_0} \cdot \frac{E}{1 + \nu} \cdot \frac{1}{\sin^2 \phi} \quad (3)$$

$$\text{where } \epsilon_{\phi} = \frac{d_{\phi} - d_0}{d_0} \cdot \frac{1}{\sin^2 \phi}$$

ϵ_{ϕ} = the strain in the surface plane in direction ϕ .

$$E = 207 \times 10^3 \text{ kg/mm}^2$$

$$\text{and } \nu = 0.27 \text{ are the constants used} \quad (4)$$

All residual stresses and strains in the EN 300 strips were found to be compressive and of the order of magnitude as published by other authors (5).

Table 1. shows that only electropolishing removed the residual strain introduced by preceding methods of surface preparation. The residual strain left after electropolishing was very small and of the same order as the limits of experimental error 34 MN/m^2 as quoted by other authors⁽³⁾.

A further EN 30B strip, heat-treated as the others, has been mechanically ground to evaluate the type and magnitude of grinding stresses. Actual D/H bit spurs are ground after braking, and hence these grinding stresses add to any residual stress already present. The grinding operation disrupted the surface to a considerable extent as has been observed for the pearlite specimens (Figure 4) ; this prevented the measurement of the elastic strain. The peak to background intensity of the diffraction profile after grinding was reduced to near zero, i.e. the peak merged with the background radiation.

The evaluation of the stress σ_ϕ from the X-ray values of interplanar strain ϵ_ϕ using Young's Modulus E and Poisson's Ratio ν , is open to criticism. The justification for this procedure, i.e., the assumption that the strain as determined by X-ray diffraction had a 1 : 1 correlation with the average strain as measured by conventional techniques, is found in the next section.

5.2 Correlation between Electrical Resistance Strain Gauge Data and X-ray Diffraction Strain Data

The four point bending stress attachment, incorporated in the modified goniometer [section 4.3] was used to set up tension in the surface of a 25-30E specimen while subject to irradiation by X-rays (Figure 5). Between the two inner points of the bending unit, an instrument rectangle of 30 x 15 mm is available, to which a 36 Ω type electrical resistance strain gauge was attached. The area next to the 36 Ω type strain gauge is still part of the instrument rectangle and is used to

obtain X-ray diffraction data. On this basis it was assumed that the strain values recorded by the SR 4 type strain gauge and those derived from X-ray diffraction are two measurements of the same strain. These strain values may differ as the SR 4 type strain gauge measured the average strain of the specimen, while the X-ray method measured only the strain on the d spacings of a particular set of crystallographic planes. Hence anisotropy of the crystal lattice may cause a variation in strain with crystallographic direction, which can become apparent as a different strain value when measured from the unique set of planes used in diffraction. As is shown next, this proved not to be the case for the specific set of crystallographic planes selected.

For hardened EN 30B specimens the (211) d spacing was selected, which diffracted at approximately $2\theta_{\mu} = 155^{\circ}$, when using chromium K_{α} radiation; this is well within the required $140^{\circ} \leq 2\theta \leq 160^{\circ}$ range.

SR 4 type strain gauges were attached to the four EN 30B strips used for experimentation in the previous section and these strips were stressed progressively in tension. At every increment the strain on the specimen was measured by a Huggenbery strainmeter. X-ray diffraction strain values were also recorded at every strain increment. The relationship between the strain gauge measurements and the X-ray diffraction strain values are presented in Figure 6.

It should be noted that the strain measurements from the SR 4 type strain gauges yielded values, which are indicative of the externally applied strain only. The residual strain present was not measured by the SR 4 type strain gauge, and this strain level is actually the arbitrary zero reference from which the externally induced tensile strains were measured. In contrast with this, the X-ray strain measurements at any stage represent the total strain level in the specimen, measuring the sum of the induced residual strain and the strain induced by the externally applied stress. Any residual strain present

before applying the external tension was revealed as a discrepancy in magnitude of the d spacing of the (211) planes which subtend different angles to the specimen surface.

The increments of strain between two levels of applied tension measured by either technique are both a measure of the same change in tension. Therefore it was necessary to determine a correlation between these two types of strain measurement.

Table II presents the data from which the graphs in Figure 6 were constructed. The relationships between the strains as measured by X-ray and strain gauge type strain gauges, depicted by these graphs, are linear and have slopes of 45° , i.e., for each increment of applied external tension, the X-ray- and strain gauge-measured strains vary by an identical amount. This indicates that the correlation between the two types of measurement is not only linear, but has a one to one correspondence, i.e., the proportionality constant necessary to equate the two strain measurements is unity.

The $\Sigma\epsilon$ values quoted in Table II, represent the total strain in the specimen surface introduced by the preparation technique and correlated with the intercepts of the linear graphs of Figure 6 with the ordinates. These values are the sum of the residual strains present in the specimen before applying an external strain.

The range in the values of $\Sigma\epsilon$ for the given external strain level applied to a specimen is maximal for specimen 3 and equals $\Delta\epsilon = .0005$. This strain value range $\Delta\epsilon$ corresponds with a double Bragg angle range of $\Delta\theta = .07^\circ$. Hence this value limits the precision of the method. Similar magnitudes of random errors in the determination of the d spacing have been published by several investigators (5).

The distribution of X-ray strain values could be due to local inhomogeneous straining induced in the crystal structure,

but it is $\epsilon \approx 20 \times$

but it is far more probable that it is due to the surface preparation. When electropolishing, it was impossible to attain a perfectly level surface free of etch pits.

The graphical relations of Figure 6 again point out the removal of residual surface strain by electropolishing. Specimens 1, 2 and 3 have undergone a final diamond polishing finish and nital etch treatment only, while specimen 4 has undergone an extensive electropolishing treatment, which had removed the strained surface layers.

From the data in Table II, and the graphs in Figure 6, it is concluded that the strains calculated from the X-ray data relating to the (211) interplanar spacings are in close agreement with the values of average strain as measured by SR 4 type strain gages. It would appear therefore that the displacement of these (211) d spacings as measured by X-ray diffraction, is representative of the strain in the whole specimen. Therefore anisotropy and lack of homogeneity, characteristic of polycrystalline materials, do not appear to exert a significant influence on the particular (211) d-spacings selected from X-ray strain measurement.

Furthermore, this direct correlation justifies the use of mechanically determined values for Young's Modulus E and Poisson's Ratio ν in evaluating the residual stresses from residual strains as measured by X-ray diffraction.

5.3 Measurements of Poisson's Ratio ν

A further indication of agreement between the SR 4 type strain gage measurements and the (211) d-spacing diffraction measurements is the close agreement in the values of Poisson's Ratio as calculated from both sets of data.

$$\nu = \frac{\epsilon_{\text{trans}}}{\epsilon_{\text{long}}} = \frac{\Delta d/d}{\Delta L/L}$$

- a) Three specimens were milled to tensile test specification from EN 30B strips and heat-treated in identical manner to the EN 30B strips as used in four point bending (Section 5.1.).

The lateral positioning of SR 4 type strain gauges on the tensile specimens yielded stress-strain data, from which Poisson's Ratio ν could be calculated and it was found that $\nu = 0.27$.

- b) Poisson's Ratio was also calculated from (211) d spacings as measured by X-ray diffraction. To this end the relationship between d_1 and d_2 from equation (2a) was used, substituting $d_1 = d_0 (1 - \frac{\nu \sigma_\phi}{E})$:

$$d_\phi = \left(\frac{1 + \nu}{E} \cdot d_0 \cdot \sigma_\phi \right) \sin^2 \Psi + d_0 \left(1 - \frac{\nu \sigma_\phi}{E} \right) \quad (11)$$

- where d_ϕ = interplanar spacing d of (211) planes at angle to the specimen surface
 d_0 = unstressed interplanar spacing d of (211) planes
 ν = Poisson's Ratio
 E = Young's Modulus = 0.2×10^{11} MN/m²
 σ_ϕ = unidirectional stress on the surface at angle ϕ to a reference line on surface

The interplanar spacing d of the set of (211) planes is calculated from X-ray diffraction data obtained at various levels of externally applied bending stress. The graphical relationship between d and $\sin^2 \Psi$ yields a set of linear graphs (Figure 7) which have a unique point of intersection. The value of $\sin^2 \Psi$ at this point of intersection is related to the value of Poisson's Ratio only and the unstressed d spacing is the unstressed d spacing d_0 . This angle Ψ is a unique angle, shown such that the negative component of stress transverse to the direction of applied compression the positive strain in the direction of stress.

Therefore Poisson's Ratio could be calculated from equation (14)

When $d_{\psi} = d_{\sigma}$, equation (14) yields the following relation

$$\sin^2 \psi = \frac{v}{1+v}$$

From Figure 7, $\sin^2 \psi = \frac{v}{1+v} = 0.22$ Hence $v = 0.28$

The two values of Poisson's Ratio as calculated in Section (a) and (b) respectively $v = 0.27$ and $v = 0.28$, agree very well with the average engineering value quoted for steels, which is $v = 0.29$. This agreement also enhances the acceptability of the (211) d spacing as a gauge length representative of the whole FN 30B specimen.

5.4 Measurement of Stress Constant K of the (211) d-spacing for Hardened FN 30B Steel

From equation (8) the stress constant K of the (211) d-spacing for hardened FN 30B steel can be calculated. Consider two stresses $(\sigma_{\phi})_1$ and $(\sigma_{\phi})_2$ of different magnitudes acting in the same direction. The corresponding strains produced are ϵ_1 and ϵ_2 , with respective diffraction angles $(\theta_1, \theta_{\psi})_1$ and $(\theta_1, \theta_{\psi})_2$.

Now substituting equation (3)

$$\epsilon_{\phi} = \frac{d_{\phi} - d_{\phi_0}}{d_{\phi_0}} = \frac{1}{\sin^2 \psi}$$

in (20) yields

$$\sigma_{\phi} = \epsilon_{\phi} \cdot \frac{E}{1+v} \quad (15)$$

Hence using equation (8)

$$K = \frac{(\sigma_{\phi})_1}{(2\theta_1 - 2\theta_{\psi})_1} = \frac{(\sigma_{\phi})_2}{(2\theta_1 - 2\theta_{\psi})_2} = \frac{(\sigma_{\phi})_1 - (\sigma_{\phi})_2}{(2\theta_1 - 2\theta_{\psi})_1 - (2\theta_1 - 2\theta_{\psi})_2}$$

Substituting σ_{ϕ} of equation (15), this yields

$$K = \frac{(\epsilon_1 - \epsilon_2) \cdot \frac{E}{1+v}}{(2\theta_1 - 2\theta_{\psi})_1 - (2\theta_1 - 2\theta_{\psi})_2} \quad (16)$$

Using the data in Table II, we may calculate K using equation (16) by substituting values for strain caused by an 4 type tension gauge

$$K = 1217 \text{ MN/m}^{-2} \cdot 10^6 \theta$$

$$\text{Hence } K = 1217$$

Using mechanical values of $E = 20.64 \times 10^4 \text{ MN/m}^2$ and $\nu = 0.29$, the stress factor K is calculated, without using the data obtained by the 3H 4 type strain gauges.

Substituting σ_4 of equation (15) into equation (8) yields the following relation for stress factor K

$$K = \frac{\sigma_4}{\epsilon} = \frac{E}{1 + \nu} \cdot \frac{1}{\sin^2 \psi} \quad (17)$$

This equation yields a value for K , the stress factor, without using X-ray data of the diffraction peak shift

$$K = 12.4 \text{ MN/m}^2 / .0108$$

Considering these two values of K it is seen that the precision of K from experimental and theoretical calculations is $\pm 5\%$, which is within the experimental X-ray diffraction error $\Delta 2\theta$. This is expected from the previous correlation between 3H 4 type strain gauge measurements and X-ray diffraction strain measurements.

5.5 Summary

1. The position of the (311) diffraction peak or the peak shift in 3H steel specimens, using chromium K_α radiation, is used to determine the residual stress removal by mechanical and electrolytic polishing of non-homogeneous plastic deformation which causes diffraction peak broadening. The same polishing technique also removes grinding, machining and other plastic stresses in the surface of hardened 3H 100 steel specimens.
2. The calibration of the X-ray strain gage of the (311) d-spacing of hardened 3H 100 steel and 3H 4 type electrical resistance strain gauge data show a correlation coefficient of unity for these two types of strain measurement. Therefore the measurement of the elastic strain on the

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1. }

a re-arrangement of the carbide particles under the influence of the surface tension forces of the binder ⁽¹⁸⁾.

The structure of a WC-Co compact is dependent upon the actual cobalt content. It was concluded from the data of several authors that for compacts with a low cobalt content, i.e., 3 volume per cent cobalt, the carbides form a continuous skeleton, while for compacts with a high cobalt content, i.e., 20 - 25 volume per cent, the structure is a cobalt-based matrix with more or less large carbide clusters embedded in it ⁽¹⁸⁾.

The structure of a 10 volume per cent cobalt-tungsten carbide compact, as used for D/H bits, is thought to consist of interlocking skeletons of tungsten carbide and cobalt ⁽¹⁸⁾.

The strength of WC-Co compacts exceeds that of the pure cobalt binder phase many times, and this is thought to be due to several factors, namely:

- a) The severe restraint exerted by the carbide particles on relative sliding of the binder
- b) The presence of residual stresses explained by the difference in coefficients of thermal expansion of both components
- c) The fact that some tungsten carbide remains in solution in cobalt at room temperature, which results in solution hardening.

A fourth point suggested to contribute to the strength of the binder, is the allotropic transformation of cobalt from a face centered cubic to a close packed hexagonal structure with its associated volume change ⁽¹⁸⁾. This point is very doubtful, as a large proportion of all the cobalt in the matrix at room temperature is in the high temperature f.c.c. form, even though it is thermodynamically unstable. The cobalt allotropic transformation is sluggish, due to the very low free energy change associated

with it, and hence the presence of F.C.C. cobalt at room temperature is very feasible. The presence of cobalt binder as a F.C.C. structure at room temperature is generally used as an industrial criterion of a satisfactory sintering technique.

The coefficients of thermal expansion of cobalt and tungsten carbide are in the ratio of three to one. The cobalt therefore will contract more than the tungsten carbide upon cooling; this sets up a tension in the cobalt and compression in the tungsten carbide grains. A mechanism explaining how these cooling stresses occur was proposed by Schilling⁽¹⁹⁾ and is outlined below.

As a result of the differential thermal shrinkage of the constituents of the compact, two mechanisms of contraction may take place. When cooling starts, the tungsten carbide grains can still move freely with respect to each other. Hence the cobalt binder, which contracts at a faster rate, can draw the tungsten carbide grains closer together. The overall shrinkage of the compact is consequently greater than that of the individual tungsten carbide grains. This mechanism can only operate as long as the tungsten carbide grains are free to move in the cobalt matrix. After the grains have made contact with each other, the compact will shrink further at the same rate as the tungsten carbide grains. The cobalt matrix is now bound in the WC grains, and hence cannot contract freely. This interaction sets up hydrostatic tensile stresses in the cobalt binder and hydrostatic compressive stresses in the tungsten carbide grains.

The final operations in the manufacture of WC-Co hard metal compacts are several mechanical treatments to achieve a satisfactory surface finish. These include tumbling, sand blasting, glass bead blasting, and vacuum blasting. These processes place the outer skin of the compact under compression, for more severely than the compression of WC grains due to thermal

contraction. Measurements of these strains are reported in the next section and yield an indication of the severity of each surface treatment operation.

6.2 Measurement of Strain in WC Grains Introduced by Surface Finishing Operations on Sintered WC-Co Compacts

The radiation selected for the diffraction of sintered 10 per cent cobalt-WC specimens was cobalt K_{α} , with an iron K_{β} radiation filter. The (hkl) reflection in the desired range $140^{\circ} \leq 2\theta \leq 160^{\circ}$ for strain measurement is that of the (202) planes of WC; their peak position is approximately $2\theta_P = 153^{\circ}$.

To measure the strain in the surface WC grains of a hard metal compact as introduced by various finishing operations, diffraction measurements were made of the as-sintered and surface treated compacts. To measure the actual strain present, no polishing or other metallographic surface preparation was applied. On the majority of the samples the surface even under different conditions of preparation is sufficiently smooth to permit substantial X-ray diffraction measurements to be made.

The data in Table III indicates the residual strain levels in the surface WC grains representative of a maximum depth of 5 μ m. The strain level of the sintered compact is negligible, unlike the remaining surface treated specimens. These had been subjected to surface preparation operations, like grit blasting, vapor blasting, tumbling, mechanical grinding and vibratory tumbling and exhibited very high levels of surface distortion. Liquidjet grinding proved to be the surface treatment which introduces minimum strain in the WC grains in the compact.

Some measurement of the severity of grit blasting and polishing with diamond paste were made also by French (22,23).

His parameter for measuring the X-ray strain was the (212) d spacing using chromium K_{α} radiation.

The limits of precision of the elastic stress measurement are again a function of the strain measurement and are $\pm 69 \text{ MN/m}^2$ for the tungsten carbide collections.

All data on the vapour blasted specimens indicate that, despite the differences in preliminary steps of preparation, the common final operation of vapour blasting was so severe that it established its own strain level eliminating any evidence of the prior level of deformation.

It has to be noted that the values of elastic strain quoted in Table III are representative only of the tungsten carbide grains. No data are included relating to the elastic straining of the cobalt binder. Therefore, according to Schminke's model of contraction events during cooling, the strain measured in the tungsten carbide grains is only a part of the total strain imposed on the hard metal compact. This was in agreement with the findings of Kochanovskaya, Keane and Marek⁽²¹⁾, who estimated that the X-ray method yielded results which were 50 per cent less than those calculated from other techniques, which measured the combined strain in WC grains and cobalt binder. Both groups of investigators attributed this discrepancy to the presence of the cobalt matrix.

This explanation is not wholly satisfactory, considering the value three investigators used for Young's Modulus when calculating the stresses. The value used in their calculations $E = 0.46 \times 10^6 \text{ N/mm}^2$ is somewhat lower than that used by Smith⁽²⁰⁾ in Table III. This value taken from Smith's data $E = 0.71 \times 10^6 \text{ N/mm}^2$ is 35 per cent greater than that used by Kochanovskaya and co-workers. D.E. French^(22,23) determined the X-ray value for Young's Modulus from measurements of the (212) d spacing using chromium K_{α} radiation, and found $E = 0.72 \times 10^6 \text{ N/mm}^2$, in full agreement with the value he quoted as reliable⁽²⁰⁾. Hence

the 40 per cent discrepancy between X-ray stress value, and stress values from other techniques, as observed by Kochanovska and co-workers, could be due to an unfortunate selection of the value of the modulus of elasticity E .

The diffraction peak shape for most of the sintered hard metal compacts was investigated for low and high angles of diffraction. The comparison of these peaks with those of an unstressed specimen has yielded information on the inhomogeneous straining and particle size which resulted in the phenomenon of peak-broadening.

To evaluate this inhomogeneous straining the diffraction peaks of atomic planes with Miller indices (001), (100), (101), (201), and (112) were investigated. The first three planes diffracted in the angular range $35^\circ < 2\theta < 60^\circ$ and in unstressed condition had a narrow base with a very high ratio of peak to background intensity. Diffraction of the last two planes (100), (112) occurred within the range $100^\circ < 2\theta < 120^\circ$, and even in the unstressed condition these peaks were relatively broad at the base, with lower ratios of peak to background intensity. The profiles of the diffraction peaks are required to calculate the magnitude of the inhomogeneous straining and the diffracting crystal size.

All reflection traces were recorded on the diffractometer chart, both for a forward and a reverse scan. The interpolation of the forward and reverse traverses was assumed to be the most reliable representation of a particular diffraction peak, as it eliminated the inertial lag of the recording unit. These interpolated peaks are represented in Figures 8, 9, 10, 11 and 12.

The diffracting crystal size of the 50 grains is dependent upon the deformation of the surface, or imposed stress

will fragment $\sqrt{> 30}$.

will fragment the WC particles into a mosaic structure with smaller areas of crystal perfection. Only crystallites smaller than 10 μm will give rise to peak broadening.

In all diffraction peak profile analyses, the actual profiles are assumed to follow either a Gaussian or Cauchy distribution curve. These profiles are defined as follows :

$$\begin{aligned} \text{Gauss profile} & : I_0 = I_o \exp(-k^2 x^2) \\ \text{Cauchy profile} & : I_C = I_o / (1 + k^2 x^2) \end{aligned}$$

This assumption constitutes an inherent weakness of these calculations as the shape of the experimental diffraction peak profile is generally intermediate between these two. In the next calculation it is assumed that both the non-homogeneous plastic deformation and the mosaic particle size independently give rise to peak broadening.

Stokes (18) expressed the diffraction peak profiles for broadened lines and reference lines of annealed samples as a Fourier series to obtain line width values for deformation and particle size only. In this project the K_{α_1} doublet, as presented by the K_{α} profiles in Figures 8 to 12, was separated by the graphical method of Kachinger (19). After this separation the breadths of the α_1 peaks for the reference specimen of annealed WC powder and the sintered compacts were calculated as the quotient of the area, A_{α_1} , under the α_1 peak (which has been corrected for background intensity) and the maximum intensity, I_{α_1} , of the α_1 peak. Next the integral breadth, β , of a broadened peak, i.e. the difference in peak breadth between the broadened peak and the reference specimen peak, is calculated from the following relationships:

$$\begin{aligned} \text{Gauss} & : \beta = B - b \\ \text{Cauchy} & : \beta = B - b \end{aligned}$$

where β = integral peak breadth
 B = peak breadth of a specimen with peak broadening
 b = peak breadth of annealed WC powder

If the diffraction peak profiles are assumed to be of the Cauchy type intensity distribution, then the following relation⁽²⁶⁾ has been derived

$$\left(\frac{\beta \cos \theta}{\lambda}\right)^2 = \left(\frac{1}{D}\right)^2 + \frac{16 \epsilon^2 \sin^2 \theta}{\lambda^2} \quad (18)$$

where D = size of diffracting crystalite

ϵ = a measure of strain

λ = wavelength in μm

Table IV. contains the values for integral breadth, β , particle size, D , and strain, ϵ , as derived from the diffraction peak profiles and equation (18).

Using the Rachinger α_1 , α_2 doublet separation construction the subsequent calculation of D and ϵ is given in Appendix B. Values for $\left(\frac{1}{D}\right)^2$ and ϵ^2 are obtained from the intercept and slope respectively of the linear relationship between $\left(\frac{\beta \cos \theta}{\lambda}\right)^2$ and $\frac{16 \sin^2 \theta}{\lambda^2}$ as presented in Figure 13. It has been possible to calculate D and ϵ only for the tumbled, electrolytically ground and grit blasted sintered compacts. The (201) and (112) diffraction peak profiles of the other specimens are not easily distinguishable from the background radiation, due to the extremely low ratio of peak to background intensity. Hence only the β values for the (001), (100) and (101) diffraction peaks were obtained for these compacts, and this lack of data prohibits the construction of the linear relationship of $\left(\frac{\beta \cos \theta}{\lambda}\right)^2$ versus $\frac{16 \sin^2 \theta}{\lambda^2}$.

Values of ϵ in Table IV. indicate that the electrolytically ground and tumbled compacts have the lowest value for non-homogeneous plastic strain. Finishing operations based on physical erosion, e.g. grinding and surface blasting by several media, introduce a far greater degree of plastic deformation of the WC grains in the sintered compact surface.

The surface roughness of the specimens can also contribute to the reduction in ratio of peak to background intensity and this is discussed in the next section. However, surface roughness measurements across the surfaces of the specimens, of which the strain data are quoted in Tables III. and IV. indicated similar roughness profiles for all specimens. Therefore, we may attribute the reduction of the intensity ratio to the surface strain alone.

The three low angle reflections of the fresh tungsten carbide powder (Figures 8,9 and 10) have well defined narrow peaks with very high ratios of peak to background intensity. This powder is almost completely strain free, as confirmed by the fact that the high angle (201) and (112) reflections show well resolved α_1 and α_2 peaks. (See Figures 11 and 12). The slightest strain would destroy this resolution and merge the particular α_1 and α_2 peaks into the usual compound α peak. Hence the tungsten carbide powder data can be used as a reference, representing material with virtually zero strain.

The diffraction peaks of the sintered only compacts show even better definition than the fresh powder peaks, having a narrower base and higher ratio of peak to background intensity. This is most probably due to grain growth of the tungsten carbide particles during the sintering process; large particles will always yield sharper diffraction lines than small particles even of similar crystal perfection. From values of B in Table IV., compared with those of the reference W₂C powder, it is concluded that the sintering process does not introduce strain on the surface layers of the compacts.

The diffraction peaks of all the other compacts, which have undergone one of the surface treatments, as outlined in Table IV., showed increased broadening of the peak and also a reduced ratio of peak to background intensity.

The difference in surface distortion between the virtually stress free sintered only compacts and the compacts subjected to various surface treatments is clearly shown by the diffraction profiles in Figures 11 and 12. All the surface treated compacts exhibit diffraction profiles for the (201) and (112) crystal planes with a composite K_{α} diffraction peak, in contrast with the well resolved K_{α_1} and K_{α_2} peaks of the sintered only compact. Besides this phenomenon a marked drop occurs in maximum peak intensity for the surface treated specimens. These high angle diffraction peaks have limited use, as a small strain tends to affect the profiles drastically, preventing the obtaining of any information on the relative difference in surface distortion between the surface treated compacts. This phenomenon is explained by Bragg's Law (equation 1), which shows that the relationship between interplanar spacing d and diffraction angle θ is a sinusoidal function, i.e., for a small variation in d , the diffraction angle will show maximum variation for crystallographic planes having large angle reflections. The low angle reflections of crystallographic planes as shown in Figures 8, 9 and 10 do indicate the difference in diffraction peak profile and to some extent the difference in distortion of the surface treated compacts. It is evident that the largest amount of distortion occurred in the ground and vapor blasted compacts. The ratio of peak to background intensity for their diffraction profiles is reduced to a value of two to one, which indicates extremely severe distortion of the surface layer.

The data quoted in Table III, for the same compacts indicate similarly large levels of residual stress for all the surface treated compacts, except for the electrolytically ground compacts. The diffraction profile distortion data in Table IV, do show in fact that the distortion is still present in the electrolytically ground compacts and that the distortion is not removed with the mechanical surface treatments. It is also evident that the distortion is not removed by the electrolytic treatment, i.e., the ground and vapor blasted compacts have a lower level of surface distortion than the electrolytically ground and vapor blasted compacts.

6.3 Surface Roughness Introduced by Surface Preparation of Sintered WC-Co Compacts

The successful measurement of residual strain by X-ray diffraction is dependent upon the condition of the surface of the specimen.

A high level of surface roughness indicates the presence of a large number of high peaks and deep valleys in the surface of the compact. This physical phenomenon causes a wide scatter of the reflected X-ray beam, which diffraction from the crystallographic planes in the surface. As the scatter increases with increasing surface roughness, the diffraction peak maximum intensity decreases until it merges with the background radiation. At a certain degree of surface roughness no satisfactory strain measurements can be made due to the large error involved in locating the peak position of a very small diffraction peak profile.

Nevertheless it is important to know these levels of surface roughness, as the surfaces will comprise numerous stress raisers introduced by abrasion or other surface finishing processes. The presence of these stress raisers may lead to early fatigue failure and hence reduce the service life of the compact. Microscopic examination of the hard metal surface in contact with the bronze metal may reduce the life of the bronze/sinter compact bond, by early failure of the WC grains adjacent to the bond.

Surface finish of the specimens for X-ray diffraction is reported in Table V. It was found that it was impossible to eliminate the plastic deformation introduced by the surface preparation process and hence some level of surface roughness was inevitable. It is therefore suggested that the surface roughness level be controlled by mechanical polishing and grinding by finishing operations.

The physical profile of a surface is measured by a surface roughness tester, which yields values for surface roughness in terms of the number of peaks and troughs on the surface and their height or depth with respect to an intermediate reference line.

The surfaces of identical sintered compacts were abraded by blasting them with a variety of media for different periods of time.

Table V. includes the values of X-ray strain measurement on these particular specimens. These data show a level of surface roughness for Samples 5, 6 and 7 which is so high that no strain data could be calculated. The amount of material removed from these three specimens was also far greater than that of the first four specimens; this factor and the surface roughness level are probably interrelated.

Specimens 8 and 9 represent two identical sintered WC compacts which have had the conventional surface treatment. The levels of surface strain of these two reference specimens are comparable with those of the first four specimens and the first four specimens are of a similar magnitude and also of the same order as the values noted in Table III. For the sintered reference specimens.

The only comparable specimen used by Brown (1953) was a grit blasted sintered compact which had a residual stress of $\sim 1311 \text{ kg/cm}^2$. This stress level is of the same order as those quoted in Tables V. and VI. These values are comparable but cannot be equated, because of possible differences in composition of the sintered material, in gritblasting conditions and in strain value as measured from the two different sets of crystallographic planes due to anisotropy of the hexagonal crystal lattice.

6.4 The Effect of Heat-Treatment on the Surface Stress of Sintered WC-Co Compacts

During the brazing cycle in the D/B bit fabrication the tungsten carbide compacts are heat-treated as described in Section 5.1. and this may have a beneficial effect in lowering the surface stress level.

Several chiselbit tungsten carbide compacts which had undergone normal surface treatment operations and exhibited a high level of surface stress were annealed in a hydrogen atmosphere at different temperatures. The levels of stress before and after heat-treatment were measured and are quoted in Table VI.

Specimen No. 1, in Table VI, indicates the residual stress of the compact prior to any heat treatment and may be taken as representative of the stress introduced by the normal surface treatment operation.

Increasing reduction in the stress level of the WC grains in the surface of the compacts is achieved by annealing at increasingly higher temperatures. Annealing the compacts for 1 hour at 900°C in a hydrogen atmosphere caused the stress level to drop off to very low values, i.e., ± 10 to $\pm 15 \text{ MN/m}^2$. This is only slightly higher than the limits of precision of stress-measurement by X-ray diffraction which is $\pm 25 \text{ MN/m}^2$. This startling reduction in stress was not observed by French (22) when heat-treating compacts of similar composition; the lowest stress level present in those compacts as reported by him was still $\pm 67 \text{ MN/m}^2$.

The reduction in the stress level by annealing in hydrogen prior to brazing indicates that during the brazing cycle in the fabrication of a D/B bit, the greater part of the residual stress, introduced into the compact during the normal surface finishing operations, is relieved. Surface stresses present in

the sintered compact surface are also expected to have been removed by this heat-treatment.

6.5 SUMMARY

The previous results show that different finishing operations introduce various degrees of surface damage and various levels of residual stress in the surfaces of the compacts.

1. An analysis of the (001), (100), (101), (201) and (111) x-ray diffraction peaks provides information on the relative surface distortion introduced by the various surface finishing operations.
2. Electrolytic grinding, which is an electrochemical erosive process, imparts a low level of residual stress and is not associated with any of the other surface finishing operations.
3. In the electrolytic grinding process, the high level of surface distortion is associated with the high level of surface distortion. The surface distortion is associated with the high level of surface distortion.
4. The surface distortion is associated with the high level of surface distortion. The surface distortion is associated with the high level of surface distortion.

The B/H bit analogue, as in the case of a B/H bit, possesses a high degree of symmetry, which directly influences the stress pattern produced by thermal shrinkage. Figure 14 is a diagram of the analogue, showing that it consists of four equivalent steel sectors, bounded by four equivalent hard metal inserts. No individual characteristic distinguishes one sector from the next and with such a symmetrical construction, each sector will have a stress pattern equivalent to the next. Obviously it is only necessary to know the stress pattern in one sector to construct the stress pattern in the whole analogue.

Consider the top right hand sector BCD in Figure 14; close examination of the geometry reveals the existence of further symmetry elements within the sector. The line bisecting angle α at C, i.e. the bisector CM, is obviously the two-fold axis of symmetry of the sector. The geometrical position of any point P(x,y) in the sector is symmetrical with the point P(y,x) in the sector. This means that the strain measurements taken in the area α' CM of the sector will yield a stress pattern in this area, which is symmetrical with the stress pattern generated in the opposite area of α' CM of the sector. Hence measurements made in area α' CM can be cross-checked by measurements in area α' CM. Upon cooling the body of the analogue contracts about its geometric center, which lies upon the axis CC (Figure 14).

Considering solely the surface plane of the analogue, it will contract equally about point C (Figure 14), but the hard metal inserts will contract to a far lesser extent about their respective geometric centers and hence stresses are generated at the points of contact between the hard metal inserts.

The variation of stress in the analogue is shown in Figure 15, which is a diagram of the stress pattern in the analogue.

cobalt compacts, as used in 'Down-the-Hole' drill bits, are described and discussed in Section 6.2. Any stresses introduced prior to the brazing operation are reduced to very low levels by the annealing operation implicit in the brazing cycle (Section 6.4., Table VI.).

Residual stresses introduced in the surface of the steel section of the D/H bit crown prior to brazing are also removed during brazing, as the EN 30B steel is austenitised during this process.

Post brazing grinding of the crown surface introduces considerable stresses in the surface, which are extremely difficult to assess quantitatively but experimental work described earlier has revealed that the residual stresses associated with the preparation of the hardened steel surface by emery paper and diamond polishing are 414 MN/m^2 in compression (Table I. Section 5.1.).

Grinding stresses in the hard metal compacts of the same magnitude have been reported by D.N. French ⁽²²⁾ at approximately -393 MN/m^2 . The reason for the unsuccessful measurement of grinding stresses in hardened EN 30B steels in these tests in contrast with French's successful measurements is that the diffraction peaks of the hard metal compacts in the as-sintered condition are clearly defined with an excellent peak to background intensity ratio, while the diffraction peaks of hardened steel, prior to any externally imposed stress, are already very broad and have low ratios of peak to background intensity. This is due to the distortion of the lattice of the steel matrix following hardening.

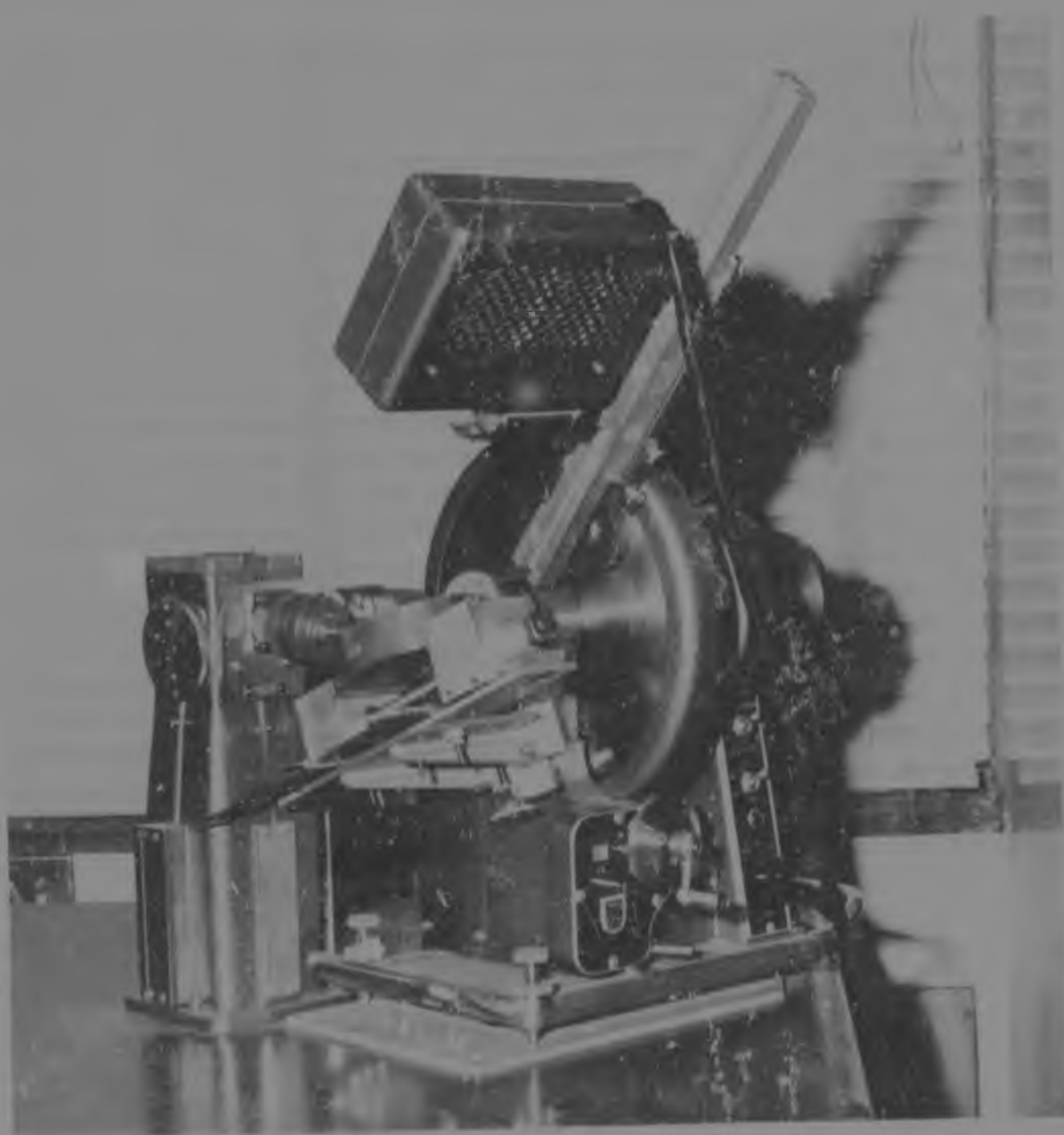
As a consequence, the grinding stresses nearly obliterate the broad martensite diffraction peaks but not the sharp, well defined tungsten carbide diffraction peaks.

In order to measure the thermal residual stress pattern in the surface of the D/H bit analogue, this specimen was prepared by the polishing technique described in Section 5.1. The analogue has four identical sectors of which one was selected and electrolytically polished. The X-ray results obtained from this sector are considered to be representative of all four sectors. The analogue situated in position for X-ray diffraction of the electropolished surface can be observed on the photograph on the next page.

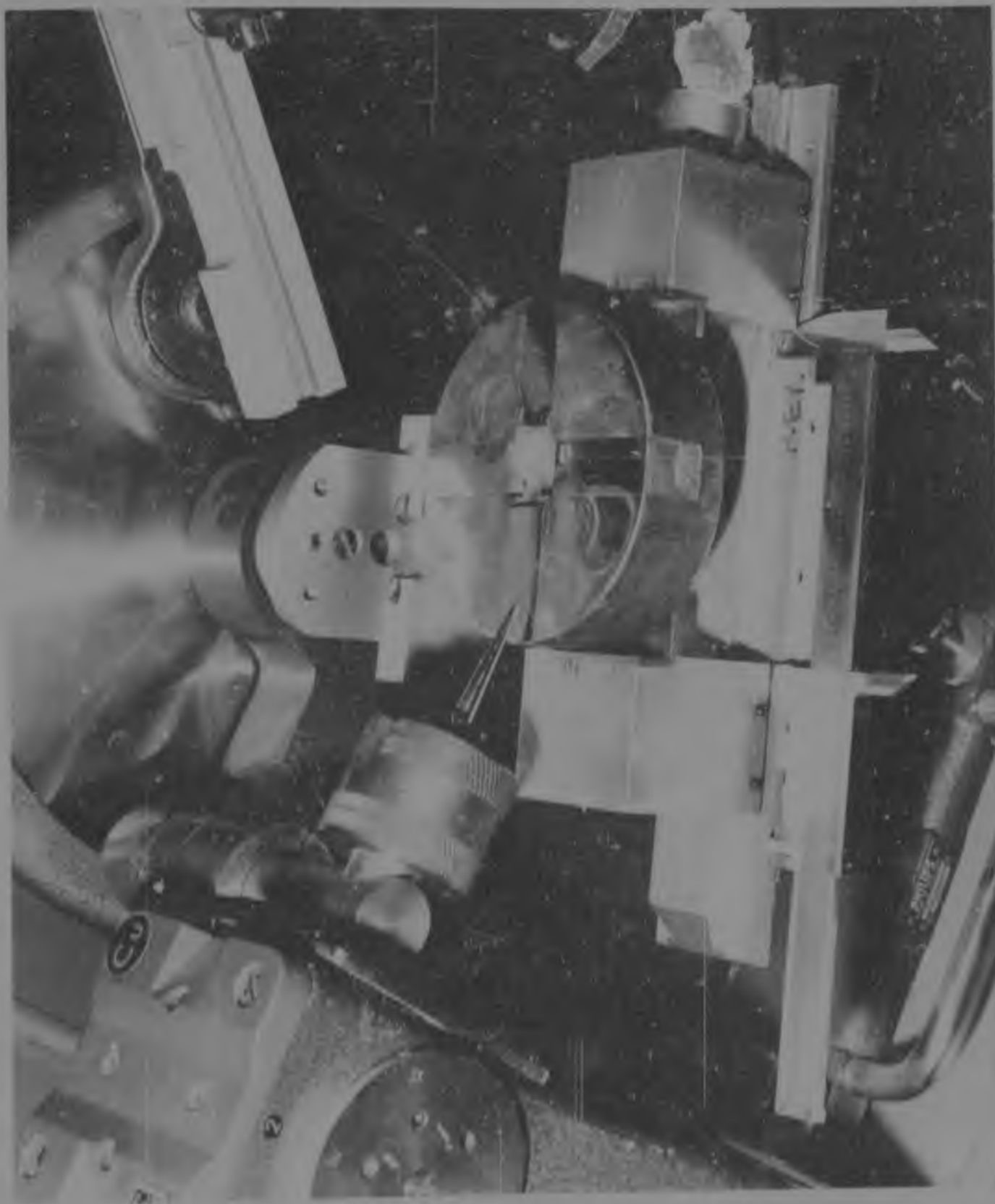
Because of the size limitations of the electrolytic polisher, three adjoining areas on the sector surface had to be polished. The boundaries between these three polished areas present problems in strain-measurement, as they exist as uneven edges of metal between the polished areas. This is in contrast with the greater centre part of the polished areas, which has a smooth bright surface. The positions of the electro-polished areas on the analogue sector surface are shown in Figure 15. From this diagram it is noted that the unevenly electro-polished areas are represented by the shaded areas A and B.

X-ray data were collected for a grid of positions in the surface of the analogue sector, as depicted in Figures 16 (a), (b) and (c). Figure 16 (a) shows the positions of the strain data taken in the x-direction; Figure 16 (b) shows the positions of the strain data taken in the 45° direction; i.e., 45° with respect to the x- and y-axes in the direction normal to the bisector of the angle (Figure 14); Figure 16 (c) shows the positions of all strain data taken in the y-direction.

The calculated strain values for each position in the respective three directions are tabulated in Tables VII, VIII and IX. A sample calculation of strain from X-ray diffraction data is given in Appendix D. Figures 17 to 25 illustrate the variation of strain in the x-direction along the y-axes and vice versa, i.e., along the x-axes.



PHOTOGRAPH I The modified X-ray unit used for the measurement of residual stress in the drill-bit analogue.



PHOTOGRAPH II. The modified X-ray unit used for the measurement of residual stresses in the drill-bit analogue - close-up view.

from Table VII. Figures 26 to 34 represent the X-ray strain-data measured in the 45° direction along the 45° traverses, i.e., data from Table VIII. Finally Figures 35 to 43 represent the X-ray strain-data measured in the y-direction along the y-traverses, i.e., data from Table IX.

All the data of these Figures 17 to 43 are plotted as strain versus distance along the respective traverse. Each calculated strain value is an average of the strain across the width (2.5 mm) of the X-ray beam and is plotted along the respective traverses at the respective co-ordinate of the centre point of the X-ray beam. Next the curves or lines of best fit were constructed to fit these data points.

Figures 17 to 25 and 35 to 43, the x- and y-strain traverses respectively, indicate the decay of strain with distance from the $x = 0$ and $y = 0$ interfaces respectively. Figures 26 to 34, representing the 45° traverses do not indicate any similar strain decay as these traverses start and end at the two stress initiating interfaces, i.e., the two interfaces fe^* and fa' .

From Figures 26 to 34 it may be seen that the strain-distance relationship for each traverse varies over a small range of strain values only. The decay of strain for the 45° traverses can only be observed when comparing the mean value of the lines of best fit for each traverse. This value decreases for each successive 45° traverse, whose distance s from point f increases. s is measured as the length of the normal from point f onto the 45° traverse, i.e., along the line fm (Figure 16 (b)).

It may be observed that the difference between the maximum and minimum values of measured strains is greater for the x- and y-traverses in Figures 17, 18, 20 and 35 to 38 than for the x- and y-traverse strain relationships in Figures 21 to 25 and 39 to 43. Therefore the precision of strain measurement

has a percentage smaller effect on the precision of the curves of best fit of Figures 17, 18, 20 and 25 than on the curves of lines of best fit of Figures 21 to 25 and 39 to 41. Consequently the curves of best fit through the data points of Figures 17, 18, 20 and 25 have a higher coefficient of correlation than the lines of best fit through the data points of the Figures 21 to 25 and 39 to 41.

This decrease in correlation between the data and the curves of lines of best fit is due to the fact that the curves of lines of best fit are not straight lines in the x, y plane but are curved surfaces, $z = f(x, y)$, in the x, y, z space.

It is impossible to obtain a straight line in the x, y, z space if the data points are not in a straight line in the x, y plane. The data points in Figures 17, 18, 20 and 25 are in a straight line in the x, y plane and the curves of lines of best fit are straight lines in the x, y, z space. The data points in Figures 21 to 25 and 39 to 41 are not in a straight line in the x, y plane and the curves of lines of best fit are curved surfaces in the x, y, z space. The curves of lines of best fit in Figures 17, 18, 20 and 25 are straight lines in the x, y, z space and the curves of lines of best fit in Figures 21 to 25 and 39 to 41 are curved surfaces in the x, y, z space.

The precision of the curves of lines of best fit in Figures 17, 18, 20 and 25 is higher than the precision of the curves of lines of best fit in Figures 21 to 25 and 39 to 41. This is due to the fact that the curves of lines of best fit in Figures 17, 18, 20 and 25 are straight lines in the x, y, z space and the curves of lines of best fit in Figures 21 to 25 and 39 to 41 are curved surfaces in the x, y, z space. The curves of lines of best fit in Figures 17, 18, 20 and 25 are straight lines in the x, y, z space and the curves of lines of best fit in Figures 21 to 25 and 39 to 41 are curved surfaces in the x, y, z space.

It may be seen from the above that the curves of lines of best fit in Figures 17, 18, 20 and 25 are straight lines in the x, y, z space and the curves of lines of best fit in Figures 21 to 25 and 39 to 41 are curved surfaces in the x, y, z space. The curves of lines of best fit in Figures 17, 18, 20 and 25 are straight lines in the x, y, z space and the curves of lines of best fit in Figures 21 to 25 and 39 to 41 are curved surfaces in the x, y, z space.

the higher degree of precision and hence are the more important for the following calculations of the residual stress pattern.

7.2 Derivation of the Residual Stress Pattern in the Analogue Surface From X-ray Strain Data

Because of experimental difficulties in locating the area to be irradiated, the three grids of ϵ_x , ϵ_y and ϵ_{45} strain data do not superimpose. For any given point P on the surface it is required to know strain data in three directions to permit evaluation of the principal stresses and their direction. It was therefore essential to derive a method which validly interpolated strain data in three directions for any point P on the surface. The graphical method used is discussed in Appendix D and its application to the strain data of Tables VII, VIII and IX yields values for the strains in directions x , y and 45° , i.e., ϵ_x , ϵ_{45} and ϵ_y . For each point P(x,y) in the surface of the sector. These strain values have been determined from two separate graphical interpolations of the experimental X-ray data for the three grids. These two separate interpolations of the same data make it possible to determine the strains in the areas of the sector for which one or the other has no equivalent values. To elucidate this matter it is necessary to consider the x- and y-traverses of Figures 16(a) and (b). Consider the ϵ_x x-traverse of Figure 16(a) and the ϵ_y and ϵ_{45} x-traverses of Figure 16(b) at the same positions on the sector surface. These exclude any strain values measured across the gap $13.7 \text{ mm} < x < 23 \text{ mm}$ as no experimental x-traverses were obtained in this gap, because it contains the uneven area A, which is the overlap of the two elliptical areas 1 and 2 of Figure 15.

The y-traverses of ϵ_x , ϵ_{45} and ϵ_y data do however yield strain values over this area from the curve of best fit through the data points. No actual measurements are made in the above gap, but the presence of strain values before and after the gap, warrant the calculation of the curve of best fit to cover the gap between the two sets of data points (as shown in Figures 35 to 43), yielding estimated strain data in this area.

However these y -traverses of σ_y , σ_{45} and σ_x data do not yield any strain data across the gap $13 \text{ mm} < x < 23 \text{ mm}$, corresponding to the uncracked area B, the overlap of the two electropolished areas I and J of Figure 15. The x -traverses of σ_x , σ_{45} and σ_y data however do yield strain data in this area for reasons similar to those discussed for the y -traverses across area A (Figure 15).

Hence the two parallel calculations serve to confirm one another and yield complementary data.

The calculation of the state of stress at any point $P(x,y)$ in the analogue sector was carried out using data from the σ_x , σ_{45} and σ_y loci of the x - and y -traverses and is presented in Table X and XI respectively. All strain data of the Tables X and XI are used in an algebraic evaluation of Mohr's circle of stress (see Appendix 1, Table I) which are recorded in the same tables.

Tables X and XI yield values for the principal strains, and also for maximum principal stresses, the angular rotation of the x -axis to the line of principal stress, the sum of the principal stresses and the difference of the principal stresses.

From the data of Tables X and XI the following relationships are plotted on a diagram of the sector surface in order to state the state of stress and rotation in the sector.

1. The σ_1 and σ_2 loci, which constitute the loci of points having equal values of principal stress σ_1 and σ_2 (Figures 61 to 64).
2. The isoclinics, which are the loci of points, tangential to which represent the principal direction of stress at the point of tangency (Figures 65, 66).
3. The isopiatics, which are the loci of points having equal values of $(\sigma_1 + \sigma_2)$ (Figures 67, 68).

4. The isochromes (Figures 69, 70).

1. Isobars

The isobar diagrams of Figures 62 and 64 depict the distribution of the principal stress (σ_2) across the surface of the analogous section. The maximum compressive stress values occur at the rounded ends of the edges (Figure 64) and equal 195 to 237 MN/m².

The compressive stress decays with increasing distance along the x- and y-oriented interfaces as well as with increasing distance towards the center of the steel sector surface.

As anticipated from the difference in brace widths, the principal stress σ_2 decays more rapidly along the y-axis than the x-axis. As a result, at approximately 20.3 mm along the x- and y-axis, the discrepancy between the (σ_2)_x and (σ_2)_y stresses has widened to 40 MN/m². Therefore the isobars are not completely symmetrical about the bisector in .

The distribution of the complementary principal stress σ_1 across the sector surface is presented in Figures 63 and 65. This principal stress σ_1 is also compressive in character, but is of a lower magnitude than σ_2 ; the maximum value for σ_1 being 55 to 72 MN/m², with distance from the center and along the braced interface, e.g., 0.5' and 0.2', σ_1 decreases to zero and then changes sign reaching a tensile stress of approximately 70 MN/m², moving towards the center and to within the corner, σ_1 decreases to zero.

Again the differing brace widths along the x- and y-axis affect the magnitude of the principal stress. At distances of 20.3 mm along the x- and y-interfaces the discrepancy in the magnitude of the principal stress σ_1 is approximately 15 MN/m². This value is similar to the quoted precision in relative stress measurements and hence is not significant. However a definite trend towards lower magnitude exists along

the y-axis and therefore this can be attributed to the differing braze widths.

The data in all four diagrams (Figures 61 to 64) are symmetrical about the bisector of within a sector of 10 mm radius from point γ . However, the variation in strain accommodation in the braze is reflected in a disruption of the symmetry of the isochromes about the bisector of this 10 mm radius.

The following anomalies occur: In the vicinity of $x = 12.7$ mm and $y = 12.7$ mm in Figures 63 and 64, i.e., towards the center of the steel sector, the values of σ_2 are larger than at adjacent positions closer to the brazed interfaces.

Furthermore the σ_1 values in the vicinity of the braze from $x = 22$ to 32 mm along the y-axis (Figures 61 and 62) also show discontinuities. The σ_1 stress values along the y-axis increase in magnitude as the braze interface is approached.

These differences are further highlighted by the graphs of the isochromes (Figures 69 and 70), which show a considerable divergence in the magnitude of angle α between the isochromes in the vicinity of the braze interface and the steel sector.

2. Isoclines

The isoclines in Figures 65 and 66 give additional information about the principal stresses. These diagrams show the direction in which σ_1 and σ_2 act in the surface of the steel sector. The isocline in the surface of the steel sector at point $P(x, y)$ represents the direction in which the respective principal stress acts.

At points close to point γ in the sector, the isochromes based on the principal stresses σ_1 and σ_2 have an equal magnitude and are directed along the y-axis.

act in directions which subtend angles of 45° to the x- and y-axes.

Along the x-axis, the effect of the stress initiated at the insert No. 2, becomes important, and the effect of the stresses generated at insert No. 1. (the y-axis) becomes negligible. Hence the principal stresses in this region act in directions parallel and normal to the brazed interface of insert No. 2, as is indicated in Figures 65 and 66. Similarly along the y-axis the principal stresses rotate from 45° to directions parallel and normal to the brazed interface of insert No. 1. Still further along the y-axis the direction of the principal stresses rotates away from these angles. Furthermore, in the area defined by $x = 10$ to 15 mm, $y = 10$ to 13 mm the directions in which the principal stresses act are erratic. These two anomalies are associated with those mentioned previously for the σ_2 isolines and are discussed later in this section.

3. Isopachics

Figures 67 and 68 represent the isopachics in the surface of the annealed sector. These curves show the magnitude of the sum of the principal stresses ($\sigma_1 + \sigma_2$), and indicate the areas of tension and compression in the steel sector surface. Near point 1, where the two compressive stresses generated at the top brazed joints are maximum, the steel sector is in considerable compression, ($\sigma_1 + \sigma_2 = -232 \text{ MN/m}^2$). Away from this area the compression is reduced as the stresses decay with distance from these points of initiation and the tensile stress induced parallel to the brazed interfaces becomes significant in magnitude. In some isolated areas, particularly along the y-axis, an unusual expansion is noted. In the area defined by $x = 0$ to 5 mm and $y = 27$ to 30 mm slight tension exists adjacent to the hinge gap in the steel, which increases in magnitude with increasing y .

The zero stress region extends over a considerable distance along $x'1$ and $x'2$, indicating that the tension generated parallel and adjacent to the brazed interfaces has a compressive counterpart of equal magnitude normal to these interfaces.

The effect of the different braze width is again apparent when considering stress distribution adjacent to the brazed interfaces. Along the y -axis the $(\sigma_1 + \sigma_2)$ stress values decay more quickly and eventually reach tensile values in contrast with the slower decay along the x -axis, where the $(\sigma_1 + \sigma_2)$ values only approach zero-stress. The difference in magnitude is again approximately 25 MN/m^2 . This phenomenon is of course expected as it is directly related to the difference in magnitude of principal stresses along the x - and y -axes.

Again the x - and y -symmetry of stress distribution is disrupted along the brazed interfaces, in contrast with the continuous symmetry between the interface and end of $x1$. The fact that symmetry along x is disrupted in contrast with the symmetry along y along the brazed interfaces, again points to the difference in braze width as the likely cause of disruption of symmetry of stress pattern.

4. Isoclines

Figures 69 and 70 show the positions of the isoclines in the stress pattern. There are the loci of points in the stress field which have the same directions of principal stresses. The angle α determines the position of the principal stress with respect to the x -axis. When $\alpha = 0^\circ$, bisecting the right angle 90° , is given as the case of symmetry, i.e. $\alpha = 0^\circ$. Any angle α is given by $\tan \alpha = \frac{\sigma_1 - \sigma_2}{2\tau}$. The direction of $\alpha = 0^\circ$ is given by $\tan \alpha = 0$ and a corresponding angle $\alpha = 90^\circ$ is this

case. Symmetry about the bisector fm is apparent, particularly in the area close to the unbrazed edges ci and af . However along the x-axis and y-axis, beyond $x = 15$ mm and $y = 15$ mm respectively, symmetry between these two axes is less apparent. Along the x-axis there exists a large area of $\alpha = 0^\circ$, while along the y-axis α decreases to zero and then changes even further to negative values. A consideration of the precision involved clarifies this anomaly which is associated with the equally anomalous principal stress values in this area and the different brace widths. The precision of strain measurement, $\epsilon = .0001$, determines the precision of the angle α at high and low values of stress. Near point f , in the steel sector, where both principal stresses are large and compressive, this precision is $\delta\alpha = 16^\circ$ (Appendix F). Along the y-axis at lower values of stress, this precision in the determination of α deteriorates to $\delta\alpha = 23^\circ$. Consequently most of the negative values for angle α along the y-axis are within the limits of precision of the calculation, assuming the true angle α to be zero degrees. Therefore, due to the large possible error in angle α at distances along the x- and y-axes, the isoclinics in these areas lose most of their significance.

A similar reasoning explains the erratic values of angle α at positions beyond point l along the line fm . Here the directions of the principal stresses vary widely, which may be due to the previously discussed low precision, which is associated with areas of low residual stress.

5. Isochromatics

The shear stresses associated with the stresses generated along the x- and y axes, are calculated from the principal stresses. The isochromatics in Figures 71 and 72 depict the distribution of double the maximum shear stress - $2\tau_{max} = (\sigma_1 - \sigma_2)$, in the surface of the steel sector.

angles are calculated from the isoclinics of Figures 69 and 70 similar erratic angles are present in the area beyond point 1 along the line fm. Here, near the line fm, the maximum shear stress directions deviate from the pattern observed in the remainder of the analogue sector.

All the previously discussed loci of Figures 61 to 74 have been calculated from the same experimental data and hence will show similar anomalies. The anomalies have been observed in areas 19 to 32 mm along the x- and y-axes and beyond point 1 along the bisector fm (Figure 14). At these positions in the steel sector, the observed residual stresses are far smaller than in the area close to point f, and hence the precision of relative stress measurement, being 34.5 MN/m^2 , will introduce a larger percentage error in these results. This is particularly important with regard to the calculation of the direction of the principal stresses and maximum shear stresses. The precision of measurement of angle α decreases from 16° near point f to 23° along the x- and y-axes.

Therefore the trends of the loci discussed in the area close to point f, i.e., the sector surface within radius 10 mm from point f, have the higher precision and are more significant.

Summarizing the residual stress pattern as measured in the surface of the steel sector and presented by Figures 61 to 74, the following description applies:

1. Both principal stresses are tensile and compressive near point f and similar principal stresses are present near the bonded interfaces aa' and ee', where σ_1 becomes tensile and σ_2 remains compressive but decreases in magnitude. This symmetry is disrupted at x = 10 mm and y = 10 mm due to the variation in strain accommodation by the different frame thicknesses.
2. The stress pattern is symmetric about the bisector fm.

Figure Maximum $\sigma_1 = 43.5$

The stress-strain characteristics of an element are represented by the relationship between forces applied to the nodal points and displacement of these nodal points, resulting from these forces. This relationship may be expressed by the flexibility or stiffness matrix of the element. The stiffness matrix of a nodal point is derived from the element stiffness matrices of those elements that share that nodal point. The equilibrium of the set of elements can be expressed by the following matrix equation

$$[F] = [K] \cdot [r]$$

where $[F]$ is the matrix of all nodal point force component matrices

$[K]$ is the matrix of all nodal point stiffness matrices

$[r]$ is the matrix of all nodal point displacement component matrices.

The above matrix equation is divided into a series of simultaneous equations, one for each nodal point, and these equations are then solved by an iterative technique using a computer program.

The structural idealisation of the analogue surface is presented in Figure 8⁴. Triangular elements were selected and the size of the smallest elements was determined by the thickness, 0.3 mm, of the brazed joint between the sintered carbide insert and the steel. The element size across the length of the brazed joint remains constant and a row of the same size elements was selected for the immediately adjoining sintered carbide and steel. At the location, where the thermal stresses are generated, i.e. at the braze, it is very important to have a very small element size, because the smaller the element size the more accurate the analysis is. With distance away from the braze into the steel and the sintered carbide, the element size is less critical and was increased to reduce the total number of elements in the whole structural assemblage.

This reduces the computing time required to complete the analysis, without materially affecting the accuracy of the results.

The program used to evaluate the structural analysis was adapted from an existing program (29), which computes thermal stresses from the material properties of the elements, the temperature interval of cooling and the co-ordinates of the nodal points.

The critical steps in this finite element analysis are the selection of the element size and the assumptions required to allow for the three dimensional analysis. Even though the X-ray strain measurements are of a plane stress system, these stresses had been generated by the thermal contraction of the three dimensional analogue and the positive volume change of the steel during the austenite to martensite transformation reaction. The finite element analysis is two-dimensional by selection and therefore fundamentally disregards the three-dimensional origin of the thermal stresses and corrections to the analysis were therefore made. Due to the limiting influence of the braze thickness on the selection of the smallest element size, a three-dimensional finite element analysis of the analogue was not attempted, because the computing time required is prohibitive.

Residual stresses in the analogue are only generated below 625°C, the braze solidification temperature. During the first few degrees of cooling below this temperature, the elastic limit of the braze is extremely small in magnitude and any stress will be relieved by plastic flow of the braze metal. In the finite element analysis it was assumed that the stresses generated over the first one hundred degrees centigrade below the braze solidification temperature are largely relieved by plastic flow and hence have a negligible magnitude. A temperature interval of 100°C was therefore assumed to be the temperature below which significant residual stress could result in appreciable

residual stress. Since no data of the magnitude of the elastic modulus of the matrix phase could be found, and since the diffusion effects between the base, steel and surface oxide are negligible, the internal stresses which develop in the matrix phase of the oxide scale are neglected. The magnitude of the elastic modulus of the oxide scale is taken to be $E = 3.5 \times 10^{11}$ dyn/cm². The Poisson's ratio of the oxide scale is taken to be $\nu = 0.35$. The thermal expansion coefficient of the oxide scale is taken to be $\alpha = 20 \times 10^{-6}$ /°C.

The austenite to martensite transformation of the EN308 steel has the following characteristic temperatures [30], $M_s = 290^\circ\text{C}$ and $M_{fin} = 180^\circ\text{C}$ and the volume change associated with this reaction is represented by the positive linear change of 56×10^{-4} inch/inch as determined dilatometrically by Russell and MacGregor [31].

During the above consideration it is assumed that phase transformation occurs over the temperature interval 334°C (phase transformation temperature) to room temperature.

1. Temperature interval 334°C to 534°C . This interval is considered a hypothetical point during the cooling cycle, above which temperature a negligible amount of residual stresses are generated, because of the elastic plasticity of the base metal above 534°C , just below the solidification temperature.
2. Temperature interval 334°C to 290°C . Over this temperature interval the steel is assumed to be elastic and hence the coefficient of thermal expansion $\alpha = 21 \times 10^{-6}$ /°C. The elastic modulus of the base is assumed to average $E = 3.5 \times 10^{11}$ dyn/cm², which allows the calculation of the stresses in the plate and the temperature distribution.

3. Temperature 290°C

3. Temperature interval 290°C to 170°C . Over this interval the steel increases in volume, the linear expansion is taken to be 56×10^{-4} inch/inch as discussed previously. Hence the average coefficient of thermal expansion of the steel is

$$\alpha = \frac{-56 \times 10^{-4}}{120} = -46.7 \times 10^{-6} / ^{\circ}\text{C}$$

This is the net result of the positive volume change associated with martensite formation and the negative volume change due to shrinkage upon cooling. The martensite reaction is estimated to have $M_f = 170^{\circ}\text{C}$. The elastic modulus of the blade is assumed to average

$$E_{290-170} = 1.9 \times 10^{11} \text{ dyn/cm}^2 = 33333 \text{ MN/m}^2$$

This decrease in magnitude allows for the still present, but rather uncertain effect on the magnitude of the elastic modulus.

4. Temperature interval 170°C to room temperature. Over this temperature interval the steel is completely martensitic and contracts according to the coefficient of thermal expansion for ferrite, $\alpha = 12 \times 10^{-6} / ^{\circ}\text{C}$. Over this temperature range the elastic modulus approaches its value of $E_{\text{ferrite}} = 190000 \text{ MN/m}^2$. The temperature effect is far less pronounced on the steel and clustered carbide components, as the cooling interval is considerably below their respective solidification temperatures.

Values used over the whole interval are

$$E_{\text{ferrite}} = 239500 \text{ MN/m}^2 \quad ; \quad \nu_{\text{ferrite}} = 0.29$$

$$\alpha_{\text{ferrite}} = 12 \times 10^{-6} / ^{\circ}\text{C} \quad ; \quad \nu_{\text{martensite}} = 0.30$$

$$E_{\text{martensite}} = 57000 \text{ MN/m}^2 \quad ; \quad \nu_{\text{ferrite}} = 0.28$$

$$\alpha_{\text{martensite}} = 33 \times 10^{-6} / ^{\circ}\text{C}$$

The change in the coefficient of thermal expansion of the steel is taken to be the same as that of the ferrite, and the change in the coefficient of thermal expansion of the steel is taken to be the same as that of the ferrite.

bisector, the line Cfm in Figure 14 which is a twofold axis of symmetry, each 45° sector is equivalent to the next and the finite element analysis may therefore be restricted to the one 45° sector.

The centre of the analogue surface plane is a void and is located in between the four sintered carbide inserts (Figure 14). Therefore a structural analysis of the 45° sector does not incorporate the fact that the analogue is a steel continuum in the volume below the sintered carbides and the central void. This presence of the steel continuum will influence the dimensional contraction of the central void.

It was attempted to incorporate the indirect effect of the steel continuum on the shrinkage of the surface plane by including elements in the surface sector of the central void, area Cfed (Figure 14), with material constants

$$\alpha = \alpha_{\text{steel}}, \quad \nu = \nu_{\text{steel}}, \quad E = \frac{E_{\text{steel}}}{10}$$

The latter reduced modulus of elasticity was so selected that the contraction of the steel continuum affects the central void and surrounding material in a reduced manner, i.e. a specific strain in the steel continuum would in the surface plane generate only one tenth of the stress it would generate the plane of the steel continuum.

The boundary conditions of the analysis, i.e. the nodal points, which have zero or one degree of freedom are the following.

1. Point C (Figure 14) is taken as the origin of the coordinate system and is defined to be fixed in both the x and y directions, i.e. it has zero degrees of freedom. In actual cooling events, C is the geometric centre of the surface plane about which the analogue surface contracts and hence has no degree of freedom.
2. The nodal points on the line Cdd have one degree of

two dimensionality of the analysis however. The σ_1 isobars of Figure 80, determined by finite element analysis, indicate the presence of stresses of compressive character near point F in the steel surface. These stresses change to tensile values with increasing value of α , i.e. towards the circumference of the sector. The similarity between the σ_1 isobars from this finite element analysis and the x-ray strain measurements, Figures 88 and 89, is pronounced in distribution and magnitude. The only difference in magnitude is the slightly higher level of tensile stresses near the circumference as computed by finite element analysis, i.e. $+150 \text{ MN/m}^2$ versus 70 MN/m^2 for the experimental work.

The maximum principal stress σ_1 in the sintered carbide surface is concentrated at either end of the insert, reaching values up to 200 MN/m^2 in tension.

The σ_2 isobars of Figure 81 in the steel surface are compressive in character and reach maximum values of 700 MN/m^2 at point F, but not at this point, the stresses decrease rapidly along line Fc and the reduction reaching zero-stress values at point c. The correspondence between the isobars in Figure 81 and those determined experimentally (Figures 82, 83) is less close than for the σ_1 isobars. The decrease in magnitude near point F is more pronounced in the isobars in Figure 81, which reach values in tension at point c. The isobars in Figure 82, which are plotted as contours of stress, show a similar pattern. The σ_2 isobars of Figure 81 in the sintered carbide surface are compressive with a slight concentration of stress at point F and along the line Fc. The isobars in Figure 83, which are plotted as contours of stress, show a similar pattern with a maximum value of 700 MN/m^2 at point F (Figure 83).

The maximum shear stresses from the finite element analysis are presented in Figure 84 as the τ_{max} isobars and show a distribution in the steel surface which is similar to that expected from the finite element analysis. The maximum shear stress in the steel surface is $\tau_{max} = 130 \text{ MN/m}^2$ in Figure 84 and $\tau_{max} = 90 \text{ MN/m}^2$ in Figure 85. The corresponding maximum shear stress in the

sintered carbide surface are concentrated parallel and adjacent to the brazed joint, having a value of 100 MN/m^2 . In the centre of the carbide insert, the magnitude has decreased to 50 MN/m^2 .

Figure 83 presents the directions of the maximum principal stress in the analogue surface. The second principal stress acts in a direction of 90° to the arrows in Figure 83, as per definition of this principal stress. From this diagram it appears as if the tensile σ_1 stresses act mainly in the width direction of the sintered carbide; the σ_2 stresses tend to act in a direction along the length of sintered carbide.

The overall residual stress pattern determined by finite element analysis of the analogue at room temperature, appears closely similar to the experimentally determined one.

The residual stresses in the sintered carbide, which have not been determined experimentally on the analogue surface, are found to have maximum tensile stresses near the 2 edges across the width of the sintered carbide, complemented by a concentration of compressive stresses adjacent to the brazed joint. These stresses are maximally $\sigma_1 = 700 \text{ MN/m}^2$ and $\sigma_2 = -300 \text{ MN/m}^2$. The maximum shear stresses are also concentrated adjacent to the braze over the length of the sintered carbide and have a maximum value of $\tau_{\text{max}} = 100 \text{ MN/m}^2$.

Because the finite element analysis was performed in four discreet cooling steps over separate temperature intervals, the stress pattern at the end of each step was plotted to analyse the change in stress distribution associated with each contraction or expansion over each temperature interval.

The residual stresses generated over the first temperature interval, 634°C to 534°C , were assumed to be negligible and Figure 76 and 77 present the finite element analysis of the stresses in the analogue at 290°C , the end of the temperature-

interval 534°C to 290°C , during which the steel is completely austenitic. The σ_1 and σ_2 isobars (Figures 76 and 77) are found to be distributed similarly to those in Figures 80 and 81, the magnitude of the isobars in Figures 76 and 77 are however found to be far in excess of those in Figures 80 and 81, with maximum values of $\sigma_1 = 600 \text{ MN/m}^2$ and $\sigma_2 = -1200 \text{ MN/m}^2$.

Figure 78 and 79 present the σ_1 and σ_2 isobars of the residual stress pattern from the finite element analysis of the analogue after cooling to 170°C , at which temperature almost all the austenite is transformed to martensite and as a result the steel over the temperature-interval 290°C to 170°C has a positive volume change, which first cancels the residual stress present at 290°C and at 170°C has its net resultant residual stresses. This cancellation and new stress pattern imposition is shown by the fact that the σ_1 isobars of Figure 78 have a similar distribution to the σ_2 isobars of Figure 77 and the σ_2 isobars of Figure 79 have a distribution similar to that of the σ_1 isobar of Figure 76.

For all three residual stress patterns at room temperature, 170°C and 290°C respectively, the directions in which the principal stresses act were found to be very similar and Figure 83 is valid for all three stress patterns. The only difference is, as expected, that the arrows indicate directions of principal stress σ_1 for the stress patterns at 290°C and room temperature, but indicate directions of principal stress σ_2 for the stress pattern at 170°C .

A summary of events generating the residual stresses during the cooling interval from brazing temperature to room temperature appears to be as follows:

- (i) Initially, just below brazing temperature, the differential thermal contraction between steel, braze and sintered carbide is absorbed by plastic deformation of the braze

which is / - 73

- which is extremely plastic at these temperatures. At lower temperatures down to 290°C , the austenitic steel, bronze metal and dispersed carbide contract differentially, which produces the stress distribution as illustrated in figures 76, 77 and 83. Very large stresses are so generated, greater in magnitude than those of the stress pattern at room temperature.
3. From the M_s temperature, 290°C , to 170°C the steel transforms to martensite with an associated volume increase, while the bronze and dispersed carbide continue to contract. This causes the cancellation of the stresses previously generated by the contraction of the austenite, and this is followed by the establishment of a second stress distribution which is in effect the inverse of the former, with σ_1 isobars resembling the former σ_2 isobars, etc. The magnitude of these stresses are also in excess of those present at room temperature. This stress pattern after the austenite is martensite positive volume change has a tensile state of stress near point P and a compressive state of stress near point Q and a large area of the steel sector. This is again shown in the graphs 78 and 79, where the tensile stress is far greater in magnitude than the compressive stress in the same positions within a radius of 10 millimeters from point P in the steel. Hence the state of stress, defined by the $(\sigma_1 + \sigma_2)$ isobar, is compressive in nature in character in contrast to the compressive state of stress for the stress pattern at 290°C and room temperatures.
4. From 170°C to room temperature the steel again contracts, as does the bronze and the dispersed carbide. This results initially in the cancellation of the stresses set up by the martensitic transformation, and is followed by the establishment of a third stress pattern defined by the contraction of all three materials. As a result the stress pattern at room temperature resembles that at 290°C but is of a much smaller magnitude.

7.5 Discussion

Even though the Finite element analysis does not allow for the ^{three-}dimensional origin of the residual stress in the analogue surface, the analysis results in a stress pattern closely similar to the experimentally determined stress pattern. This has the following implications and uses:

1. The computer output yielding a hypothetical residual stress pattern from finite element analysis may be considered as an approximation of the experimentally determined stress pattern in the steel surface and to evaluate the analogue surface experimental procedures.
2. The residual stress pattern in the welded carbide inserts which could not be determined using the X-ray diffraction technique, may be assumed to resemble the residual stress pattern in the welded carbide inserts from the finite element analysis.
3. The residual stress pattern in the analogue surface at various temperatures may be assumed to resemble the stress pattern obtained from the finite element analysis of the analogue differential thermal contraction at these temperatures.

During cooling from brazing temperature the behaviour of the steel matrix causes three successive residual stress patterns to be established.

Over the temperature interval 170°C to room temperature the residual stress pattern present at 170°C is cancelled by the contraction of the matrix, which opposes the volume increase from the austenite to martensite transformation which is established the residual stress pattern present at 170°C .

It is proposed that at some temperature between room temperature and 170°C the analogue surface is almost completely free from any residual stresses due to differential thermal

contraction effects between the sintered carbide compact and the steel, and that the only residual stresses present are those generated by differential thermal contraction effects between the bronze metal and the adjoining steel and the sintered carbide respectively. This latter stress pattern will be of comparatively negligible magnitude in the steel and the sintered carbide, because any differential thermal contraction is mainly absorbed by the bronze, as it has the lower modulus of elasticity and hence absorbs the greater amount of the strain generated.

It is further proposed that the residual stresses as determined for the analogous surface will be very similar to the residual stress pattern in a plane normal to the longitudinal axis of a down-the-hole drill bit intersecting the sintered carbide inserts through their upper portion below the cutting faces. The absence of rigidity will introduce a less steep stress gradient in the steel of the analogue, but the stresses at the brazed interfaces should be similar because the finite element analysis indicated that these are purely a function of the cruciform geometry, the temperature-interval of cooling and the material constants. Similarly the absence of steel stock underneath the sintered carbide inserts in the analogue will only affect the stress distribution at the circumference, but not at the brazed interfaces.

Therefore the residual stress pattern in the analogous surface directly adjacent to the brazed interfaces are thought to be very similar to the stresses in similar positions in a down-the-hole drill bit. The stresses at the circumference may differ between the analogue and the drill bit, due to more pronounced stress relief in the analogue steel by bending, which in the case of the drill bit is opposed by the rigidity of a long shaft of drill steel.

The reduction in the magnitude of the residual stress pattern with an increase in temperature above room temperature can occur in practice, when under drilling conditions the drill head heats up. If the temperature of the drill head increases considerably during drilling, the compressive nature of the residual stresses at the centre, near point r , may change to a tensile nature. This type of stress distribution is not favourable for fatigue loading conditions, which are present during drilling, and hence elevated temperatures during drilling should be avoided to prolong the working life of a drill.

7.6 Summary

1. It is anticipated that machining stresses in the steel surface of the drill crown, introduced prior to brazing, are substantially removed by the austenitisation of the EN 30B steel during brazing.
2. Grinding stresses of considerable compressive magnitude are imposed on the drill crown surface after brazing. These stresses destroy the thermal residual stress pattern in the surface layers of the drill crown. This thermal residual stress pattern is generated during the cooling period from the brazing temperature and is present in the surface layers underneath the post-brazing grinding stresses.
3. The position of the nearly vertical axis between the two steel edges, defining the brazed interfaces is a two-fold axis of symmetry in the sector surface. This may be deduced from geometric considerations and is substantiated by the X-ray strain data. Symmetry of the residual stress pattern is not complete about this line but is probably disrupted by residual stress relaxation in the braze annulus with the different braze widths. Further, the geometric centre of the analogue the precision of the measurement of the stresses is highest and hence the

1001 In this area are the more significant indicators of the stress pattern.

- a. The principal stresses in the steel are compressive and at a maximum near the centre of the analogue, the value being $\sigma_1 = -70 \text{ MN/m}^2$ and $\sigma_2 = -237 \text{ MN/m}^2$.
- b. In the steel, away from the centre of the analogue and parallel to the two brazed interfaces, the principal stress σ_1 is tensile, the maximum value being $\sigma_1 = +90 \text{ MN/m}^2$. Normal to the brazed interfaces the principal stress σ_2 remains compressive, decreasing in magnitude towards the circumference of the analogue.
- c. The principal stresses in the steel area adjacent to the two-fold axis of symmetry are parallel to this line. These directions change to 45° to the bisector, i.e. normal and parallel to the brazed interfaces, in positions further away from the axis of symmetry.
- d. The state of stress in the steel sector area varies from compression of stress type, as indicated by the $(\sigma_1 + \sigma_2)$ isopneustics.
- e. The direction of the maximum shear stress in the steel sector varies from parallel to the brazed interfaces, at the two-fold axis of symmetry, i.e. parallel with the brazed interfaces, with increasing distance from the bisector.

The maximum shear stress occurs adjacent to the brazed interfaces at positions $x = 5.4$ to 12.7 mm along the x-interface and $y = 5.4$ to 12.7 mm along the y-interface. The maximum value of shear stress obtained is

$$\tau_{\max} = 97 \text{ MN/m}^2.$$

4. A finite element analysis is used to calculate the thermal residual stresses produced from the expansion restraint and the mechanical constraints of the analogue components. This structural analysis yields a stress pattern, which is similar to the as measured stress pattern of

$$\text{error } \tau_1 / -7\% -$$

point 3.

5. The principal stresses in the surface of the hard metal inserts, balancing the residual stresses in the steel sector across the brazed interfaces, are assumed to be similar to those calculated by the finite element analysis. This assumption is made on the strength of the similarity between the stress patterns in the steel as determined from X-ray strain data and from finite element analysis.

The maximum stresses in the sintered carbide are

respectively $\sigma_1 = 200 \text{ MN/m}^2$, $\sigma_2 = -300 \text{ MN/m}^2$.

$\tau_{\max} = 100 \text{ MN/m}^2$. Stress σ_1 is concentrated at the two edges of the carbide across its width, σ_2 is concentrated parallel and adjacent to the brazed joint, $\tau_{\max}$ is also concentrated parallel and adjacent to the brazed joint.

6. The residual stress distribution in the analogue surface at elevated temperatures is thought to be similar to those calculated by finite element analysis on the strength of the correspondence between the two stress distributions; experimental and from finite element analysis, at room temperature.

8. Conclusions

1. A reliable X-ray method for residual stress measurement has been developed for EN 30B hardened steel. The method is based on the use of a specially designed X-ray diffractometer with a goniometer, using this X-ray method, residual stresses were measured over small circular areas of approximately 5.0 square millimeters which permits the application of the method to the measurement of residual stress in small areas.

2. A comparative investigation of the various surface treatment processes, which are or may be used during the final stages of production of sintered tungsten carbide compacts, indicates that mechanical grinding, tumbling and shot and vapour blasting techniques introduce very high residual stresses and considerable plastic deformation of the surface layers of the sintered compacts. A non-physical erosion finishing operation, viz., an electrolytic grinding process, was found to introduce minimal residual stress and least plastic deformation of the surface layers.

The greater proportion of all the residual stresses was found to be removed by the annealing process implicit in the brazing operation.

3. The close correspondence between the residual stress patterns in the analogue steel surface as constructed from X-ray experimental data and from finite element analysis has the following implications:
- a. The laboratory experimental technique has validated the finite element analysis of cooling events in the analogue, so that the latter may be used for further residual stress problems under similar circumstances.
 - b. The residual stress pattern at elevated temperatures may be estimated using the finite element analysis.
 - c. Residual stresses in the sintered carbide compacts may be assumed to be similar to those determined by finite element analysis.
4. The residual stress pattern resulting from grinding and brazing in the surface layers of a cruciform drill bit is complex. The extreme surface layers are in considerable compression. The magnitude and distribution of this stress is not uniform and is dependent on the geometry of the drill bit. It is important to note that the large stresses are concentrated in the surface layers and are relieved by the large stresses in the bulk of the material.

introduced during the post-brazing grinding. Underneath these grinding stresses, the residual stress pattern, generated by the differential thermal contraction, persists. An increase in operating temperature of the drill is thought to decrease the magnitude of the latter type of stress pattern, changing it to a tensile character. This could be detrimental to drill performance, as a tensile type of stress pattern is generally not beneficial under fatigue-loading conditions present during drilling.

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APPENDIX 'A'

ELECTROPOLISHING OF HARDENED EN 30B STEEL

^a / Electrolyte used

5.0 volume per cent H_2O
18.5 volume per cent $HClO_4$ 70 per cent
76.5 volume per cent $(CH_3CO)_2O$

Current density on surface of approximately 600 mm^2

3.9 - 5.4 mAmp/mm^2

APPENDIX 'B'

The Calculation of Particle Size D and Lattice Strain
of an Electrolytically Ground Sintered Compact

The K_{α} profiles of (001), (100), (101), (201) and (112) diffraction peaks are used to calculate the lattice-strain ϵ and particle size D.

The values for β are taken from Table IV. The following are the data required to evaluate graphically the Cauchy relationship.

$$\left(\frac{\beta \cos \theta}{\lambda} \right)^2 = \left(\frac{1}{D} \right)^2 + \frac{16 \epsilon^2 \sin^2 \theta}{\lambda^2} \quad (1)$$

(hkl)	$\frac{16 \sin^2 \theta}{\lambda^2}$	$\left(\frac{\beta \cos \theta}{\lambda} \right)^2$
001	49.6	.018
100	63.1	.100
101	112.4	.181
201	302.0	.233
112	388.5	.177

The linear relationship of the equation (1) is defined by the data in above table. The construction of the locus is presented as line (1) in Figure 15, which has

$$\begin{aligned} \text{slope} &= \frac{0.10}{300} = 0.00033 = \epsilon^2 \longrightarrow \epsilon = 0.0182 \\ \text{intercept} &= 0.08 \mu\text{m}^{-2} = \left(\frac{1}{D} \right)^2 \longrightarrow D = 3.5 \mu\text{m}. \end{aligned}$$

APPENDIX 'C'

Sample Calculation of Strain ϵ ϕ from X-ray Data
In Direction ϕ in a Steel Surface

X-ray intensity data measured for specimen surface angles $\psi = 0^\circ$ and $\psi = 45^\circ$ in direction ϕ are the following:

$\psi = 0^\circ$ $2\theta^\circ$	Number of Pulses from counter	Time I secs	Time II secs	Time III secs	I_{ψ}/I_0
$2\theta_1 = 154.7$	63348	2000	2520	2400	0.785
$2\theta_2 = 155.7$	58	2000	1970	1388	1.000
$2\theta_3 = 156.7$	60613	2000	2540	2538	0.744

$\psi = 45^\circ$ $2\theta^\circ$	Number of Pulses from counter	Time I secs	Time II secs	Time III secs	I_{ψ}/I_0
$2\theta_1 = 154.8$	53080	1200	1631	1210	0.757
$2\theta_2 = 155.8$	70960	1200	1218	916	1.000
$2\theta_3 = 156.8$	57636	1200	1500	1146	0.800

The above data include intensity as number of pulses from the proportional counter, received by the electronic counting circuit. The time interval (I) is the time period over which pulses are counted, given time (II) represents the number of seconds required to reach an intensity count of 6000 and 7000 pulses respectively for each angular setting $2\theta_1$, $2\theta_2$ and $2\theta_3$.

Appendix (C)

Data in Column Time (III) are the same time periods as recorded in Column Time (II), divided by the θ -dependent correction factors of Richlars and Shipley ⁽⁴⁾, which correct the absorption dependent asymmetry of the diffraction peaks.

The last Column, $I_{0/L}$ represents the intensity ratios of the $2\theta_1$ and $2\theta_2$ positions relative to the $2\theta_0$ position, which is taken to be 1.0. These intensity ratios are next used in the algebraic evaluations of the parabolic equation to obtain the peak position $2\theta_p$.

$$2\theta_p = 2\theta_1 + \frac{c}{2} \left(\frac{3a + b}{a + b} \right)$$

where for $\psi = 0^\circ$
 $a = 0.215$ $b = 0.256$ $c = 1.0^\circ \cdot 2\theta$

hence $(2\theta_p)_0 = 154.7 + \frac{1.0}{2} \left(\frac{0.645 + 0.256}{0.215 + 0.256} \right) = 155.656^\circ$

and for $\psi = 45^\circ$
 $a = 0.243$ $b = 0.200$ $c = 1.0^\circ \cdot 2\theta$

hence $(2\theta_p)_{45} = 155.847^\circ$

Calculate n_ψ from equation (3) and equation (1)

$$n_\psi = \frac{d_\psi - d_1}{d_1} = \frac{1}{\sin^2 \psi} \quad (3)$$

$$n\lambda = 2d \sin \theta \quad (1)$$

hence $d_\psi = \frac{n\lambda}{2 \sin \theta_\psi}$

and $d_1 = \frac{\lambda}{2 \sin \theta_1}$

so that $n_\psi = \frac{1/\sin \theta_\psi - 1/\sin \theta_1}{1/\sin \theta_1} = \frac{1}{\sin^2 \psi}$

for $\psi = 0^\circ$, $\sin \theta_1 = \sin 77.828^\circ = .9775$

$\psi = 45^\circ$, $\sin \theta_\psi = \sin 77.925^\circ = .9779$

$$n_\psi = \left\{ \frac{1}{\sin^2 \psi} - \frac{1}{\sin^2 \theta_1} \right\} (1/.9775) = (1.02260 - 1.02302)(.9775)$$

$$= -.00041 \quad \text{i.e., a compressive strain.}$$

APPENDIX 'D'

Construction of Strain Values in Directions $\pm 45^\circ$
and y for each point $P(x,y)$ in the Steel Sector

To obtain the residual strain data for each point $P(x,y)$, the following constructions and calculations are carried out:

1. Using the respective strain data from the curves or lines of best fit of Figures 35 to 43, the x -direction traverse of the strain data in the y -direction, ϵ_y , is constructed; i.e. from each line or curve of best fit one strain value at a specific position y_1 (distance from insert No. 2) is obtained and all these strain values are used to construct the graph of ϵ_y versus distance along the x -direction traverse, which is the line parallel to the x -axis (insert No. 2) at distance y_1 .
2. Similarly the x -direction traverse of all the strain data in the 45° direction, ϵ_{45} , from Figures 26 to 34 is constructed along the line $y = y_1$.
3. Using the x -direction traverse of ϵ_y and ϵ_{45} and the original x -direction traverse ϵ_x , the strain distribution at all points $P(x,y)$ along the x -direction is calculated.

Analogous calculations and constructions are performed for the ϵ_x and ϵ_{45} values applied for all x -direction traverse along line $y = y_1$ together with the x -direction traverse of ϵ_y values. Hence the strain distribution for all points $P(x,y)$ in the sector surface is known.

Analogous calculations and constructions are performed for the ϵ_y and ϵ_{45} values applied for all y -direction traverse along line $x = x_1$ together with the y -direction traverse of ϵ_x values. Hence the strain distribution for all points $P(x,y)$ in the sector surface is known.

At this point $n = 2$

Appendix D

At this point it is necessary to comment that the transposition of the strain data as described above introduces new errors. Besides the experimental error, due to equipment and surface preparation, the subsequent use of the curves of best fit through the transposed strain data may introduce an additional error.

Example - Construction of x-direction Traverse of y-direction Strain Data

The following construction is an example of the X-ray strain data transpositions.

Consider the $y = 1.27$ mm. traverse, i.e. the traverse of x-direction strain data (Figure 17) along the traverse $y = 1.27$ mm.

It is the aim to construct the 45° of the y-direction and 45°-direction strain data along the same $y = 1.27$ mm. traverse. To construct these two traverses, the y-direction and 45°-direction strain data at $\Delta y = 1.270$ mm. of curves in Figures 35 to 43 and Figures 26 to 34 respectively are acquired and plotted versus their respective Δx values along the $y = 1.27$ mm. traverse. Hence now for this traverse there exist three curves of strain in x, 45° and y-direction respectively versus distance in the x-direction. The data acquired from Figures 26 to 43 are quoted below.

Appendix 1b*

Δx mm	Δy mm	ϵ_y	Δx mm	Δy mm	a mm	b mm	ϵ_{45}
2.54	1.27	-.00090	0.53	1.27	1.27	0.75	-.00155
3.81	1.27	-.00100	2.33	1.27	2.54	3.30	-.00134
6.99	1.27	-.00068	0.32	1.27	5.71	8.25	-.00084
10.16	1.27	-.00056	11.33	1.27	8.90	16.00	-.00045
13.32	1.27	-.00039	16.00	1.27	10.79	19.82	-.00030
22.84	1.27	-.00038	16.70	1.27	12.70	22.88	-.00020
26.18	1.27	-.00040	27.87	1.27	17.78	33.76	-.00007
31.78	1.27	-.00038	30.60	1.27	22.56	41.92	-.00016
36.95	1.27	-.00045	38.43	1.27	26.66	51.54	-.00007

The two transposed x-traverses for ϵ_y and ϵ_{45} along $\Delta y = 1.27$ mm. can now be constructed. The y- and 45° -direction strain data, ϵ_y and ϵ_{45} , of Figures 25 to 43 and 26 to 34 respectively are transposed in an analogous manner to the x-direction traverses over the range $5.82 \leq y \leq 31.75$ mm yielding analogous transposed x-traverses.

The x-traverses for all ϵ_y and ϵ_{45} data are presented in Figures 44 to 51.

The y-traverses for all ϵ_x and ϵ_{45} data are presented in Figures 52 to 60.

APPENDIX 1C

Mohr's Circle of Strain

Ref. p. 33-35 G.B. Dieter Mechanical Metallurgy
McGraw Hill 1961

The above reference describes the graphical evaluation of principal strain ϵ_1 and ϵ_2 , shear strain γ and direction α of principal strains with respect to the x-direction from experimental values of the strains ϵ_x , ϵ_{45} and ϵ_y .

These three strain data ϵ_x , ϵ_{45} and ϵ_y completely define the Mohr circle and a computer program was written to circumvent the laborious graphical method of construction.

This computer program is based upon the following relationships

Mohr circle : Center $(\epsilon_{11}, \gamma_{11})$, radius r
Principal strains are

$$\epsilon_1 = \epsilon_{11} + r$$

$$\epsilon_2 = \epsilon_{11} - r$$

$$\epsilon_{11} = \frac{\epsilon_x + \epsilon_y}{2}$$

$$r = \sqrt{\left(\frac{\epsilon_x - \epsilon_y}{2}\right)^2 + (\gamma_{45})^2}$$

$$2\alpha = \cos^{-1} \left(\frac{\epsilon_x - \epsilon_{11}}{r} \right)$$

Angle α is measured with respect to the x-direction and is always positive except for the case

$$\epsilon_x \leq \epsilon_{45} \leq \epsilon_y, \text{ when } |\epsilon_x - \epsilon_{45}| > |\epsilon_y - \epsilon_{45}|$$

EXAMPLES

$$1. \quad e_x = -.00078, \quad e_{45} = -.00010, \quad e_y = +.00032$$

$$\text{then } x_{11} = -.00055$$

$$\text{and } r = \sqrt{\frac{(.00068)^2 + (.00022)^2}{2}} = .000505,$$

$$\text{and } \angle \alpha = \cos^{-1} \left(\frac{-.00078 + .00032}{.000505} \right) = 62.9^\circ,$$

$$\text{hence } \alpha = 31^\circ, \quad e_1 = -.00106, \quad e_2 = -.00005,$$

$$2. \quad e_x = -.00040, \quad e_{45} = 0, \quad e_y = +.00020$$

$$\text{then } x_{11} = -.00010$$

$$\text{and } r = \sqrt{\frac{(.0004)^2 + (.0002)^2}{2}} = 0.000316$$

$$\text{hence } \angle \alpha = \cos^{-1} \left(\frac{-.00040 + .00010}{.000316} \right) = 18.4^\circ,$$

$$\text{and } \alpha = -9^\circ, \quad \text{as } |e_x - e_{45}| > |e_y + e_{45}|$$

$$\text{and } e_x < e_{45} < e_y.$$

$$e_1 = -.00040, \quad e_2 = +.00020$$

APPENDIX (F)

Calculation of Error $\Delta \alpha$ Associated with Angle α For Low and High Levels of Horizontal Strain

The precision of angle α for any set of strain data is a function of the magnitude of those strain data and its precision $\delta e = .00010$.

From Appendix E the relationship between angle α and strain data x_1 , x_2 , and x_3 is

$$2\alpha = \cos^{-1} \frac{\frac{e_x - e_y}{2}}{\sqrt{\frac{(e_x - e_{45})^2 + (e_y - e_{45})^2}{2}}}$$

The maximum variation in angle 2α is

$$(2\alpha \pm \delta 2\alpha) = \cos^{-1} \left[\frac{\frac{(x_1 + x) - (x_3 - x)}{2}}{\sqrt{\frac{(x_1 - x - x_2 - x)^2 + (x_3 - x - x_2 - x)^2}{2}}} \right]$$

$$= \cos^{-1} \left[\frac{\frac{(x_1 - x_3) / 2 + x}{\sqrt{\frac{(x_1 - x_2)^2 + (x_3 - x_2)^2}{2} - \delta x(2x_1 - 4x_2 + 2x_3) + 4\delta e^2}}}{\sqrt{\frac{(x_1 - x_2)^2 + (x_3 - x_2)^2}{2} - \delta x(2x_1 - 4x_2 + 2x_3) + 4\delta e^2}} \right]$$

$$\text{where } e_x = x_1 \quad e_{45} = x_2 \quad e_y = x_3$$

1. Low Level of Strain

$$\text{e.g. } x_1 = .00010 \quad x_2 = .00006 \quad x_3 = .00022$$

$$2\alpha = 29.0^\circ \quad \alpha = 14.5^\circ$$

$$(2\alpha \pm \delta 2\alpha) = 14.5^\circ \quad \alpha \pm \delta \alpha = 7.25^\circ$$

$$\text{Hence precision } \delta \alpha = .29^\circ$$

Appendix III

2. High Level of Seepage

$$\text{R.C. } \lambda_1 = -.00110 \quad \lambda_2 = -.00748 \quad \lambda_3 = -.00122$$

$$2 \cdot \alpha = 78^\circ \quad \rightarrow \quad \alpha = 39^\circ.$$

$$(\alpha + \delta \alpha) = 46^\circ \quad \rightarrow \quad (\alpha - \delta \alpha) = 23^\circ.$$

$$\text{Revised Transmittion } \delta \alpha = 16^\circ.$$

APPENDIX 56A

Width of Braze between Hard Leather and Hard
Metal Inserts Nos. 1 and 2 of the Anafogue

Microscopic measurements, at intervals of 1.0 mm. along
the braze, yielded the following data:

1. Braze next to hard metal insert (1): 0.32 mm $\longleftrightarrow$ 0.40 mm
Average : 0.35 mm.
2. Braze next to hard metal insert (2): 0.28 mm $\longleftrightarrow$ 0.35 mm.
Average : 0.315 mm.

However, 90 per cent of the width of this braze gap lies
within the limits

$$0.24 \text{ mm.} \longleftrightarrow 0.29 \text{ mm.}$$

$$\text{Average is } 0.265 \text{ mm.}$$

TABLE 1.

DIFFRACTION DATA AND DERIVED RESIDUAL STRAIN AND STRESS
VALUES ON 1% C STEEL AFTER METALLOGRAPHIC POLISHING
AND BRINDING

Specimen Preparation	θ_{ϕ}	$\theta_{2\theta}$ p	$\sin \theta$ p	$\frac{d_h - d_i}{d_i}$	STRAIN e_{ϕ}	STRESS σ_{ϕ} MN/m ²
1						
Emery Paper Grind and Diamond Polish	0	155.176	.97663	-.00128	-.0026	-414
	45	155.877	.97790			
2						
As above	0	155.335	.97691	-.00126	-.0025	-393
	45	156.019	.97814			
3						
As above	0	155.343	.97691	-.00108	-.0022	-345
	45	155.914	.97798			
4						
As above and Electropolish	0	155.965	.97804	-.00000	-.0001*	-14*
	45	155.990	.97810		-.0002	-34

* A tensile strain of .0001 was placed on Specimen 4 to enable it to remain in position in the goniometer; with no external strain it had a tendency to slip. Hence the measured results of $e_{\phi} = -.0001$ and $\sigma_{\phi} = -14$ MN/m² are not the actual residual strain and stress.

The true residual strain and stress are also quoted in Table 1 and are

$$e_{\phi} = -.0002 \quad ; \quad \sigma_{\phi} = -34 \text{ MN/m}^2.$$

TABLE II.

A COMPARISON OF STRAIN AS MEASURED BY SR 4 STRAIN GAUGES
AND X-RAY DIFFRACTION

	ϵ_{SR4}	$\psi = 0^\circ$ $\frac{\Delta\theta}{2\theta} \frac{a}{p}$	$\psi = 45^\circ$ $\frac{\Delta\theta}{2\theta} \frac{a}{p}$	X-ray strain ϵ_x	$\Sigma \epsilon$
<u>SPECIMEN 1</u>	0	155.74	155.91	-.0022	-.0022
Emery paper	+.0006	155.73	155.82	-.0018	-.0024
grind and	+.0012	155.46	155.74	-.0012	-.0024
Diamond	+.0020	155.55	155.65	-.0005	-.0023
polish	+.0024	155.59	155.60	-.0001	-.0025
<u>SPECIMEN 2</u>	0	155.18	155.88	-.0026	-.0026
	+.0009	155.28	155.73	-.0018	-.0027
As above	+.0016	155.77	155.61	-.0009	-.0027
	+.0027	155.49	155.48	+.0001	-.0026
<u>SPECIMEN 3</u>	0	155.34	155.62	-.0025	-.0025
	+.0003	155.22	155.84	-.0023	-.0026
As above	+.0006	155.22	155.82	-.0022	-.0028
	+.0009	155.33	155.73	-.0016	-.0025
	+.0018	155.39	155.61	-.0009	-.0027
	+.0027	155.46	155.50	-.0001	-.0028
<u>SPECIMEN 4</u>	+.0001	155.97	155.99	-.0001	-.0002
	+.0006	156.02	155.91	+.0003	-.0003
As above -	+.0012	156.07	155.83	+.0008	-.0004
Electropolished	+.0018	156.14	155.78	+.0015	-.0003
	+.0026	156.24	155.68	+.0021	-.0003

ϵ_{SR4} = uniaxial strain imposed on specimen by 4-point bending and measured by SR 4 strain gauges.

X-ray strain ϵ_x = total elastic strain present in specimen as measured by X-ray diffraction.

$$\Sigma \epsilon = \epsilon_x - \epsilon_{SR4}$$

N.B.

The SR 4 resistance strain gauges and the X-ray apparatus were both aligned to measure in the direction of the applied stress, i.e., along the longitudinal axis of the 30P strip.

TABLE III.

X-RAY MEASURED ELASTIC STRAINS AND THEIR CORRESPONDING
STRESSES IN THE SURFACE OF SINTERED WC-CO COMPACTS

Specimen	e Compressive	Compression
		σ MN/m ²
1. As sintered	.0001	55
2. Tumbled	.0014	770
3. Grit blasted	.0015	825
4. Electrolytically Ground	.0004	220
5. Mechanically Ground	.0012	660
6. Vapour blasted	.0014	770
7. Alundum and Vapour blasted	.0014	770
8. Tumbled and Vapour blasted	.0014	770
9. Vibratory tumbled (1)	.0014	770
Vibratory tumbled (2)	.0017	935
10. Sintered and Gritblasted (D.N. French)		1311

The value of Young's Modulus used was obtained from
 literature (1965) as $E = 0.70 \times 10^6 \text{ MN/m}^2$

TABLE IV.

STRAIN VALUES AND PARTICLE SIZES DETERMINED FROM THE INTEGRAL BROADENING β

OF DIFFRACTION PEAK PROFILES

Specimen description	DIFFRACTION PEAKS												ϵ
	(100)		(101)		(201)		(112)				D μm		
	β	B	β	B	β	B	β	B					
1. WC powder	.22	.22	.20	.20	.35	.40							
2. Sintered only	.13	.15	.15	.15	.29	.36							
3. Electrolytically ground	.30	.08	.41	.10	.29	.78	.90	.50	.50	3.5		.0182	
4. Tumbled	.41	.10	.44	.22	.45	.83	1.14	.74	.74	3.8		.0277	
5. Gritblasted	.56	.34	.59	.37	.68	1.38	1.03			2.2		.0535	
6. Ground	.32	.10	.67	.45	.73	.53							
7. Vapour blasted	.32	.10	.67	.45	.73	.53							
8. Aluminium and vapour blasted	.42	.10	.67	.45	.73	.53							
9. Tumbled and vapour blasted	.32	.10	.67	.45	.73	.53							

Area under $K\alpha_1$ diffraction peakMax. intensity of $K\alpha_1$ peak $\beta_{\text{broad}} = \beta_{(hkl)}$ broadened - $\beta_{(hkl)}$ WC powder

N.B. Specimens 2 to 9 inclusive are sintered compacts

TABLE V.
X-RAY MEASUREMENT ON THE SURFACE OF WC-Co SINTERED COMPACTS HAVING DIFFERENT LEVELS OF
SURFACE ROUGHNESS

Abrasive	Pressure kg/m ²	Process Time (seconds)	Surface Roughness		Thickness		Compressive Strain e	Compressive σ MN/m ²
			Before μ m	After μ m	Before mm	After mm		
1. 400 mesh bauxite	586	30	.305	.886	11.02	11.01	.0023	1256
2. 400 mesh bauxite	586	30	.432	.889	11.02	11.01	.0020	1092
3. 400 mesh bauxite	586	30	.432	.889	11.01	10.99	.0022	1201
4. 45/75 Vibro-peads	586	30	.432	.559	11.01	11.01	.0019	1037
5. 80/20 Al ₂ O ₃ ide	586	30	.432	1.905	10.99	10.95	Measurement not possible because of extreme rough- ness of surface.	
6. 80/20 Al ₂ O ₃ ide	586	30	.406	2.032	11.00	10.93		
7. 80/20 Al ₂ O ₃ ide	586	30	.432	2.032	11.00	10.81		
8. Standard Hard Metals blasting (South Africa)							.0021	1147
9. Standard Hard Metals blasting (South Africa)							.0021	1147
10. D.K. French. Sintered and Gritblasted								1314

VALUES FOR STRESS WERE CALCULATED AS FOR TABLE III.

TABLE VI.

EFFECT OF HEAT TREATMENT ON THE LEVEL OF RESIDUAL STRESS
IN SINTERED CARBIDE SURFACES

All specimens were acquired from Boart and Hard Metals (S.A.) Limited and had been processed with the normal surface blasting technique.

<u>No.</u>	<u>Heat Treatment</u>			<u>Compressive</u>	<u>Compressive</u>
	<u>Temperature</u>	<u>Time</u>	<u>Atmosphere</u>	<u>Strain</u> e	<u>Stress σ</u> MN / m ²
1	-	-	-	.0026	1430
2	520°C	1 hour	H ₂	.0011	605
3	520°C	1 hour	H ₂	.0014	770
4	700°C	1 hour	H ₂	.0005	275
5	700°C	1 hour	H ₂	.0005	275
6	900°C	1 hour	H ₂	.0003	165
7	900°C	1 hour	H ₂	.0002	110

Stress σ was calculated as per values in Table III.

TABLE VII.

RESIDUAL STRAINS IN X-DIRECTION ACROSS THE STEEL SECTOR
SURFACE CALCULATED FROM X-RAY DIFFRACTION PEAK SHIFTS

The relevant X-ray data are quoted as the double diffraction angles $2\theta_p$ for the two angles of specimen surface rotation

$$\psi = 0^\circ \quad \text{and} \quad \psi = 45^\circ$$

dx = distance from point f along insert (2).

dy = distance from f along insert (1).

POSITION		$\psi = 0^\circ$	$\psi = 45^\circ$	$(2\theta_0 - 2\theta_{45})$	STRAIN
dx mm	dy mm	$2\theta_p^\circ$	$2\theta_p^\circ$	$\Delta 2\theta_p^\circ$	e_x
0.64	1.27	155.597	155.814	-0.217	-.00085
1.27	1.27	155.606	155.860	-0.254	-.00099
1.91	1.27	155.544	155.793	-0.249	-.00097
3.28	1.27	155.642	155.743	-0.101	-.00039
4.45	1.27	155.663	155.741	-0.078	-.00030
5.82	1.27	155.666	155.778	-0.112	-.00044
8.36	1.27	155.725	155.790	-0.065	-.00025
10.90	1.27	155.759	155.739	+0.020	+.00008
13.44	1.27	155.789	155.691	+0.098	+.00038
15.98	1.27	155.839	155.714	+0.125	+.00049
24.14	1.27	155.774	155.660	+0.114	+.00045
26.00	1.27	155.714	155.653	+0.061	+.00024
27.96	1.27	155.750	155.654	+0.096	+.00037
29.82	1.27	155.764	155.659	+0.105	+.00041
31.75	1.27	155.738	155.682	+0.056	+.00022
33.64	1.27	155.727	155.630	+0.097	+.00038
35.58	1.27	155.726	155.647	+0.079	+.00031
1.27	3.81	155.654	155.871	-0.217	-.00085
1.91	3.81	155.733	155.815	-0.082	-.00032
3.28	3.81	155.633	155.774	-0.141	-.00055
4.45	3.81	155.673	155.734	-0.061	-.00024
5.82	3.81	155.720	155.730	-0.010	-.00004
8.36	3.81	155.719	155.673	+0.046	+.00018
10.90	3.81	155.808	155.693	+0.115	+.00045
13.44	3.81	155.747	155.684	+0.063	+.00025
15.98	3.81	155.781	155.724	+0.057	+.00022
24.14	3.81	155.735	155.700	+0.035	+.00014
26.00	3.81	155.704	155.659	+0.045	+.00021
27.96	3.81	155.711	155.660	+0.051	+.00020
29.82	3.81	155.725	155.710	+0.015	+.00003
31.75	3.81	155.726	155.661	+0.065	+.00025
33.64	3.81	155.677	155.657	+0.020	+.00008
35.58	3.81	155.698	155.646	+0.052	+.00020

continued

TABLE VII.
(continued)

POSITION		$\psi = 0^\circ$	$\psi = 45^\circ$	$(2\theta_0 - 2\theta_{45})$	STRAIN
dx mm	dy mm	$2\theta_p^\circ$	$2\theta_p^\circ$	$\Delta 2\theta_p^\circ$	e_x
1.91	5.82	155.811	155.851	-0.04	-.00016
3.28	5.82	155.662	155.776	-0.116	-.00045
4.45	5.82	155.709	155.830	-0.121	-.00047
5.82	5.82	155.721	155.777	-0.056	-.00022
8.36	5.82	155.698	155.752	-0.054	-.00021
10.90	5.82	155.733	155.736	-0.003	-.00001
13.44	5.82	155.780	155.750	+0.030	+.00012
15.98	5.82	155.798	155.700	+0.098	+.00038
0.64	8.00	155.852	155.952	-0.100	-.00039
1.91	8.00	155.693	155.810	-0.117	-.00046
3.28	8.00	155.677	155.768	-0.091	-.00036
4.45	8.00	155.653	155.853	-0.200	-.00078
5.82	8.00	155.712	155.790	-0.078	-.00030
8.36	8.00	155.724	155.772	-0.048	-.00019
10.90	8.00	155.743	155.771	-0.028	-.00011
13.44	8.00	155.725	155.726	-0.001	.00000
15.98	8.00	155.801	155.689	+0.112	+.00044
21.60	8.00	155.827	155.731	+0.096	+.00037
24.15	8.00	155.747	155.652	+0.095	+.00037
26.68	8.00	155.783	155.723	+0.060	+.00023
29.20	8.00	155.782	155.740	+0.042	+.00016
31.70	8.00	155.751	155.720	+0.031	+.00012
34.30	8.00	155.725	155.746	-0.021	-.00008
0.64	9.53	155.754	155.860	-0.106	-.00041
1.91	9.53	155.751	155.849	-0.098	-.00038
3.28	9.53	155.718	155.894	-0.176	-.00069
4.45	9.53	155.724	155.890	-0.166	-.00065
5.82	9.53	155.808	155.851	-0.043	-.00017
8.36	9.53	155.763	155.819	-0.056	-.00022
10.90	9.53	155.710	155.842	-0.132	-.00052
13.44	9.53	155.683	155.799	-0.116	-.00045
15.98	9.53	155.776	155.838	-0.062	-.00024

continued

TABLE VII.
(continued)

POSITION		$\psi = 0^\circ$	$\psi = 45^\circ$	$(2\theta_0 - 2\theta_{45})$	STRAIN
dx mm	dy mm	$2\theta_p^\circ$	$2\theta_p^\circ$	$\Delta 2\theta_p^\circ$	e_x
0.64	12.70	155.753	155.772	-0.019	-.00007
1.91	12.70	155.769	155.852	-0.083	-.00032
3.28	12.70	155.741	155.846	-0.105	-.00041
4.45	12.70	155.767	155.787	-0.020	-.00008
5.82	12.70	155.764	155.780	-0.016	-.00006
8.36	12.70	155.757	155.793	-0.036	-.00014
10.90	12.70	155.776	155.763	+0.013	+.00005
13.44	12.70	155.693	155.829	-0.136	-.00053
14.60	12.70	155.757	155.811	-0.054	-.00021
0.64	22.86	155.733	155.754	-0.021	-.00008
1.91	22.86	155.735	155.766	-0.031	-.00012
3.28	22.86	155.728	155.748	-0.020	-.00008
5.82	22.86	155.728	155.733	-0.005	-.00002
8.36	22.86	155.737	155.746	-0.009	-.00004
10.90	22.86	155.738	155.771	-0.033	-.00013
13.44	22.86	155.745	155.728	+0.017	+.00007
15.24	22.86	155.753	155.720	+0.033	+.00013
0.64	27.94	155.689	155.754	-0.065	-.00025
3.28	27.94	155.689	155.802	-0.113	-.00044
5.82	27.94	155.748	155.745	+0.003	+.00001
8.36	27.94	155.710	155.743	+0.027	+.00011
10.90	27.94	155.723	155.739	-0.016	-.00006
13.44	27.94	155.709	155.711	-0.002	-.00001
15.24	27.94	155.720	155.684	+0.036	+.00014
0.64	31.75	155.730	155.783	-0.053	-.00021
1.91	31.75	155.725	155.753	-0.028	-.00011
3.28	31.75	155.708	155.815	-0.107	-.00042
5.82	31.75	155.729	155.794	-0.065	-.00025
8.36	31.75	155.718	155.763	-0.045	-.00018
10.90	31.75	155.700	155.735	-0.035	-.00014
13.44	31.75	155.724	155.741	-0.017	-.00007

TABLE VIII.

RESIDUAL STRAINS IN THE 45° DIRECTION ACROSS THE STEEL
SECTOR SURFACE CALCULATED FROM X-RAY DIFFRACTION PEAK SHIFTS

The relevant X-ray data are quoted as the double diffraction angle $2\theta_p$ for the two angles of specimen surface rotation

$$\psi = 0^\circ \text{ and } \psi = 45^\circ$$

b - distance along 45° traverse, measured from the y-axis

s - distance of 45° traverse line from point f, measured along the normal fm

P O S I T I O N				$\psi = 0^\circ$	$\psi = 45^\circ$	$\Delta 2\theta_p$	STRAIN
s mm	b mm	dx mm	dy mm	$2\theta_p^\circ$	$2\theta_p^\circ$	$(2\theta_0 - 2\theta_{45})$	e_{45}
1.27	0	0	1.90	155.520	155.965	-0.455	-.00174
1.27	1.27	0.95	0.95	155.604	155.972	-0.368	-.00144
1.27	2.54	1.90	0	155.535	155.999	-0.464	-.00181
2.54	0	0	3.81	155.593	155.883	-0.290	-.00113
2.54	1.27	0.95	2.86	155.602	155.825	-0.223	-.00087
2.54	2.54	1.90	1.90	155.610	155.940	-0.330	-.00129
2.54	3.81	2.86	0.95	155.592	155.949	-0.357	-.00139
2.54	5.08	3.81	0	155.597	155.874	-0.277	-.00108
5.71	0.64	0.51	7.62	155.637	155.804	-0.167	-.00065
5.71	3.28	2.28	5.84	155.623	155.853	-0.230	-.00090
5.71	5.72	4.06	4.06	155.670	155.929	-0.259	-.00101
5.71	8.26	5.84	2.28	155.670	155.875	-0.205	-.00080
5.71	10.80	7.62	0.51	155.610	155.821	-0.211	-.00082
8.90	0	0	12.70	155.724	155.868	-0.144	-.00058
8.90	2.54	1.78	10.63	155.664	155.786	-0.142	-.00055
8.90	5.08	3.56	8.90	155.721	155.797	-0.071	-.00028
8.90	7.62	5.34	7.11	155.710	155.858	-0.122	-.00048
8.90	10.16	7.11	5.30	155.656	155.854	-0.198	-.00077
8.90	12.70	8.90	3.56	155.678	155.868	-0.170	-.00060
8.90	15.24	10.63	1.78	155.692	155.871	-0.131	-.00051
8.90	17.78	12.70	0	155.712	155.795	-0.083	-.00032

continued . . .

TABLE VIII.
(continued)

P O S I T I O N				$\psi = 0^\circ$	$\psi = 45^\circ$	$\Delta 2\theta^\circ$	STRAIN
s mm	b mm	dx mm	dy mm	$2\theta^\circ$ _P	$2\theta^\circ$ _P	$(2\theta^\circ_0 - 2\theta^\circ_{45})$	ϵ_{45}
10.79	0	0	15.87	155.763	155.715	+0.048	+0.00019
10.79	2.54	1.78	14.35	155.696	155.760	-0.064	-0.00025
10.79	5.08	3.56	12.58	155.693	155.790	-0.097	-0.00038
10.79	7.62	5.34	10.80	155.732	155.796	-0.064	-0.00025
10.79	12.70	8.90	8.90	155.700	155.816	-0.116	-0.00045
10.79	15.24	10.80	5.39	155.679	155.794	-0.115	-0.00045
10.79	17.78	12.58	3.56	155.693	155.782	-0.089	-0.00035
10.79	20.32	14.35	1.78	155.672	155.786	-0.114	-0.00045
10.79	21.59	15.87	0	155.729	155.757	-0.028	-0.00011
12.70	0.64	0.64	18.40	155.712	155.747	-0.035	-0.00017
12.70	3.18	2.28	16.63	155.738	155.697	+0.041	+0.00010
12.70	5.72	4.06	14.86	155.758	155.742	+0.016	+0.00006
12.70	8.26	5.84	13.08	155.721	155.778	-0.057	-0.00022
12.70	13.33	9.40	9.40	155.720	155.831	-0.111	-0.00043
12.70	18.40	13.08	5.84	155.700	155.812	-0.112	-0.00044
12.70	20.96	14.86	4.06	155.710	155.802	-0.092	-0.00036
12.70	23.48	16.63	2.28	155.760	155.811	-0.051	-0.00020
12.70	24.79	18.40	0.64	155.729	155.756	-0.027	-0.00011
17.78	0	0	25.40	155.683	155.689	-0.006	-0.00002
17.78	2.54	1.78	23.38	155.769	155.691	+0.078	+0.00030
17.78	5.08	3.56	21.60	155.734	155.723	+0.011	+0.00004
17.78	7.62	5.34	19.80	155.696	155.745	-0.049	-0.00019
17.78	12.70	8.90	16.12	155.749	155.751	-0.002	-0.00001
17.78	17.78	12.57	12.57	155.693	155.760	-0.067	-0.00026
17.78	22.86	16.12	8.90	155.717	155.773	-0.056	-0.00022
17.78	27.94	19.80	5.34	155.742	155.712	+0.030	+0.00012
17.78	30.48	21.60	3.56	155.760	155.718	+0.042	+0.00016
17.78	33.02	23.38	1.78	155.723	155.700	+0.023	+0.00009
17.78	35.56	25.40	0	155.700	155.708	-0.008	-0.00003

continued

TABLE VIII.
(continued)

P O S I T I O N				$\psi = 0^\circ$	$\psi = 45^\circ$	$\Delta 2\theta_p^\circ$	STRAIN
s	b	dx	dy	$2\theta_p^\circ$	$2\theta_p^\circ$	$(2\theta_0 - 2\theta_{45})$	e_{45}
mm	mm	mm	mm				
22.56	0	0	31.76	155.687	155.569	+0.118	+0.00046
22.56	2.54	1.78	30.46	155.659	155.603	+0.056	+0.00022
22.56	5.08	3.56	28.70	155.665	155.634	+0.031	+0.00012
22.56	7.62	5.34	26.92	155.731	155.634	+0.097	+0.00038
22.56	12.70	8.90	23.00	155.749	155.681	+0.068	+0.00027
22.56	17.78	12.57	19.30	155.774	155.704	+0.070	+0.00027
22.56	24.13	17.00	14.86	155.713	155.726	-0.013	-0.00005
22.56	30.48	21.46	10.29	155.753	155.732	+0.021	+0.00008
22.56	35.56	24.79	6.73	155.734	155.725	-0.009	+0.00004
22.56	38.10	26.92	4.95	155.709	155.714	-0.005	-0.00002
22.56	40.64	28.70	3.18	155.752	155.678	+0.074	+0.00029
22.56	43.18	30.46	1.40	155.713	155.673	+0.040	+0.00016
22.56	45.10	31.76	0	155.690	155.669	+0.021	+0.00008
26.66	0	0	38.10	155.739	155.648	+0.091	+0.00036
26.66	2.54	1.78	36.10	155.686	155.640	+0.046	+0.00018
26.66	5.08	3.56	34.18	155.700	155.682	+0.018	+0.00007
26.66	7.62	5.34	32.40	155.726	155.666	+0.060	+0.00023
26.66	12.70	8.90	28.80	155.683	155.753	-0.070	-0.00027
26.66	17.78	12.57	25.16	155.744	155.730	+0.014	+0.00005
26.66	27.94	19.80	18.02	155.737	155.738	-0.001	0
26.66	35.56	25.16	12.51	155.743	155.741	+0.002	+0.00001
26.66	40.64	28.80	8.90	155.733	155.721	+0.012	+0.00005
26.66	45.72	32.40	5.34	155.713	155.748	-0.035	-0.00014
26.66	48.20	34.18	3.56	155.739	155.748	-0.009	-0.00004
26.66	50.80	36.10	1.78	155.695	155.756	-0.061	-0.00024
26.66	53.34	38.10	0	155.683	155.678	+0.005	+0.00002

TABLE IX.

RESIDUAL STRAINS IN THE Y-DIRECTION ACROSS THE STEEL SECTOR
SURFACE, CALCULATED FROM X-RAY DIFFRACTION PEAK-SHIFTS

The relevant X-ray data are quoted as the double diffraction angle $2\theta_p$ for the two angles of specimen surface rotation

$$\psi = 0^\circ \quad \text{and} \quad \psi = 45^\circ$$

dx = distance from point f along insert (2)

dy = distance from f along insert (1)

POSITION		$\psi = 0^\circ$	$\psi = 45^\circ$	$\Delta 2\theta_p$	STRAIN
dx	dy	$2\theta_p$	$2\theta_p$	$(2\theta_0 - 2\theta_{45})$	e_y
mm	mm				
2.54	0.64	155.571	155.847	-0.276	-.00108
2.54	1.27	155.602	155.819	-0.217	-.00085
2.54	2.54	155.635	155.794	-0.159	-.00062
2.54	3.81	155.652	155.784	-0.132	-.00052
2.54	5.35	155.741	155.768	-0.027	-.00011
2.54	8.89	155.767	155.733	+0.034	+0.00013
2.54	11.43	155.783	155.736	+0.047	+0.00018
2.54	13.97	155.753	155.712	+0.041	+0.00016
2.54	24.79	155.738	155.686	+0.052	+0.00020
2.54	27.30	155.722	155.688	+0.034	+0.00013
2.54	29.82	155.702	155.625	+0.077	+0.00030
2.54	32.40	155.737	155.673	+0.064	+0.00025
3.18	0.64	155.638	155.902	-0.264	-.00103
3.18	1.27	155.634	155.908	-0.274	-.00107
3.18	2.54	155.616	155.839	-0.223	-.00087
3.18	3.81	155.690	155.864	-0.174	-.00068
3.18	5.08	155.714	155.852	-0.138	-.00054
3.18	6.35	155.746	155.822	-0.076	-.00030
3.18	8.89	155.703	155.763	0	0
3.18	11.43	155.793	155.742	+0.051	+0.00020
3.18	13.97	155.770	155.725	+0.045	+0.00018
3.18	16.50	155.793	155.732	+0.061	+0.00025
3.18	27.30	155.784	155.717	+0.067	+0.00026
3.18	32.40	155.766	155.672	+0.094	+0.00037
3.18	34.90	155.743	155.645	+0.098	+0.00038
3.18	37.45	155.767	155.646	+0.121	+0.00047

continued ...

TABLE IX.
(continued)

POSITION		$\psi = 0^\circ$	$\psi = 45^\circ$	$\Delta 2\theta^\circ$	STRAIN
dx mm	dy mm	$2\theta^\circ$ _P	$2\theta^\circ$ _P	$(2\theta^\circ_0 - 2\theta^\circ_{45})$	e_y
6.98	0.64	155.690	155.828	-0.138	-.00054
6.98	1.27	155.651	155.852	-0.201	-.00078
6.98	2.54	155.660	155.832	-0.172	-.00067
6.98	3.81	155.706	155.785	-0.079	-.00031
6.98	5.08	155.677	155.807	-0.130	-.00051
6.98	6.35	155.671	155.847	-0.176	-.00069
6.98	7.62	155.710	155.775	-0.065	-.00025
6.98	8.89	155.727	155.788	-0.061	-.00024
6.98	11.43	155.782	155.741	+0.041	+.00016
6.98	13.97	155.790	155.728	+0.062	+.00024
6.98	16.51	155.789	155.680	+0.109	+.00043
6.98	24.79	155.787	155.660	+0.127	+.00050
6.98	26.00	155.7	155.677	+0.120	+.00047
6.98	27.30	155.800	155.673	+0.127	+.00050
6.98	29.82	155.853	155.717	+0.136	+.00053
6.98	32.40	155.778	155.672	+0.106	+.00041
6.98	34.90	155.751	155.647	+0.104	+.00041
6.98	37.45	155.775	155.680	+0.095	+.00037
10.16	0.64	155.711	155.818	-0.107	-.00042
10.16	1.27	155.686	155.867	-0.181	-.00071
10.16	1.91	155.723	155.854	-0.131	-.00051
10.16	2.54	155.717	155.840	-0.123	-.00048
10.16	3.81	155.733	155.863	-0.130	-.00051
10.16	5.08	155.762	155.833	-0.071	-.00028
10.16	6.35	155.742	155.846	-0.104	-.00041
10.16	7.62	155.753	155.815	-0.062	-.00024
10.16	8.89	155.751	155.779	-0.048	-.00019
10.16	10.16	155.763	155.720	+0.043	+.00017
10.16	11.43	155.774	155.751	+0.023	+.00009
10.16	12.70	155.800	155.750	+0.050	+.00020
10.16	13.97	155.807	155.743	+0.064	+.00025
10.16	17.78	155.822	155.708	+0.114	+.00044

continued

TABLE IX.
(continued)

POSITION		$\psi = 0^\circ$	$\psi = 45^\circ$	$\Delta 2\theta_p$	STRAIN
dx mm	dy mm	$2\theta_p^\circ$	$2\theta_p^\circ$	$(2\theta_0 - 2\theta_{45})$	e_y
13.32	0.64	155.777	155.866	-0.089	-.00035
13.32	1.27	155.766	155.875	-0.109	-.00043
13.32	2.54	155.772	155.867	-0.095	-.00037
13.32	3.81	155.722	155.838	-0.116	-.00045
13.32	6.35	155.754	155.876	-0.122	-.00048
13.32	8.89	155.681	155.835	-0.154	-.00050
13.32	11.43	155.708	155.820	-0.112	-.00044
13.32	13.97	155.740	155.750	+0.010	+.00004
13.32	16.51	155.825	155.772	+0.053	+.00021
22.86	0.64	155.731	155.843	-0.112	-.00044
22.86	1.91	155.711	155.794	-0.083	-.00032
22.86	3.28	155.714	155.795	-0.081	-.00032
22.86	5.82	155.708	155.755	-0.047	-.00018
22.86	8.36	155.718	155.748	-0.030	-.00012
22.86	10.90	155.723	155.738	-0.015	-.00006
22.86	13.44	155.731	155.721	+0.005	+.00002
22.86	15.98	155.705	155.732	-0.027	-.00011
26.18	0.64	155.655	155.805	-0.150	-.00059
26.18	1.91	155.710	155.768	-0.058	-.00023
26.18	3.28	155.671	155.773	-0.102	-.00040
26.18	5.82	155.702	155.745	-0.043	-.00017
26.18	8.36	155.688	155.721	-0.033	-.00013
26.18	10.90	155.681	155.743	-0.062	-.00024
26.18	13.44	155.703	155.730	-0.027	-.00011
26.18	15.98	155.719	155.720	-0.001	0
26.18	18.40	155.697	155.702	-0.005	-.00002
31.76	0.64	155.705	155.775	-0.070	-.00027
31.76	1.91	155.673	155.800	-0.127	-.00050
31.76	3.28	155.706	155.763	-0.057	-.00022
31.76	5.82	155.694	155.744	-0.050	-.00020
31.76	8.36	155.720	155.715	+0.005	+.00002
31.76	10.90	155.731	155.760	-0.029	-.00011
31.76	13.44	155.721	155.720	+0.001	0
31.76	15.98	155.685	155.666	+0.019	+.00007

continued

TABLE IX.
(continued)

POSITION		$\psi = 0^\circ$	$\psi = 45^\circ$	$\Delta 2\theta_p$ ($2\theta_0 - 2\theta_{45}$)	STRAIN e_y
dx mm	dy mm	$2\theta_p$	$2\theta_p$		
34.90	0.64	155.667	155.802	-0.135	-.00053
34.90	1.91	155.691	155.791	-0.100	-.00039
34.90	3.28	155.655	155.836	-0.171	-.00067
34.90	5.52	155.678	155.761	-0.083	-.00032
34.90	8.36	155.698	155.741	-0.043	-.00017
34.90	10.90	155.678	155.682	-0.004	-.00002

TABLE 3.

PRINCIPAL STRESSES AND ASSOCIATED STRESS LOGS, DERIVED FROM MOHR'S CIRCLE OF STRAIN
CALCULATIONS USING X-RAY DATA

Δx	Δy	e_x	e_{45}	e_y	e_z	e_1	σ_1^p	σ_1	σ_2	$\sigma_1 + \sigma_2$	$\sigma_1 - \sigma_2$
1.27	1.27	-.00080	-.00147	-.00070	-.00162	-.00003	43	-5	-237	-242	232
2.54	1.27	-.00068	-.00132	-.00050	-.00123	-.00025	51	-40	-198	-235	158
3.81	1.27	-.00055	-.00116	-.00041	-.00122	-.00034	61	-55	-197	-252	142
5.08	1.27	-.00043	-.00102	-.00029	-.00109	-.00023	61	-37	-176	-213	139
6.35	1.27	-.00030	-.00088	-.00075	-.00094	-.00010	61	-16	-151	-167	133
7.62	1.27	-.00018	-.00076	-.00066	-.00084	0	63	0	-135	-135	135
10.16	1.27	+.00007	-.00053	-.00052	-.00066	.00021	66	34	-106	-72	140
12.70	1.27	+.00011	-.00037	-.00042	-.00059	.00043	70	69	-87	-18	156
15.24	1.27	+.00047	-.00029	-.00040	-.00049	.00055	74	89	-77	12	164
20.32	1.27	+.00043	-.00006	-.00040	-.00041	.00045	85	72	-66	6	138
25.40	1.27	+.00018	+.00010	-.00040	-.00042	.00030	98	64	-68	-4	142
30.48	1.27	+.00033	+.00016	-.00040	-.00043	.00038	104	61	-72	-11	135
35.56	1.27	+.00028	-.00002	-.00040	-.00040	.00024	67	45	-61	-19	109
2.54	3.81	-.00050	-.00106	-.00049	-.00111	.00007	45	+11	-179	-168	190
3.81	3.81	-.00033	-.00097	-.00046	-.00104	-.00004	54	-2	-161	-163	159
5.08	3.81	-.00020	-.00088	-.00044	-.00093	.00019	56	14	-150	-136	164
6.35	3.81	-.00006	-.00080	-.00059	-.00087	.00022	60	35	-140	-105	175
7.62	3.81	+.00009	-.00072	-.00053	-.00082	.00030	62	58	-132	-74	190
10.16	3.81	+.00028	-.00053	-.00047	-.00068	.00040	65	79	-109	-30	188

TABLE X.
PRINCIPAL STRESSES AND ASSOCIATED STRESS FACT, DERIVED FROM MOHR'S CIRCLE OF STRAIN
CALCULATIONS USING X-RAY DATA

$\Delta\epsilon$	$\Delta\gamma$	e_x	e_{45}	e_y	p_2	p_1	α_x	σ_1	σ_2	$\sigma_1 + \sigma_2$	$\sigma_1 - \sigma_2$
12.70	3.81	+.00028	-.00039	-.00042	-.00034	.00040	70	64	- 87	- 23	131
17.78	3.81	+.00026	-.00006	-.00034	-.00034	.00026	88	42	- 55	- 13	97
22.86	3.81	+.00024	+.00026	-.00036	-.00038	-.00032	110	52	- 61	- 9	119
27.04	3.81	+.00022	+.00020	-.00030	-.00030	.00031	111	50	- 64	- 13	113
33.02	3.81	+.00020	+.00019	-.00030	-.00040	.00030	112	48	- 64	- 16	112
8.54	5.71	-.00036	-.00089	-.00020	-.00090	.00034	41	+55	-145	- 90	200
3.81	5.71	-.00034	-.00082	-.00042	-.00082	.00006	46	10	-132	-122	142
5.08	5.71	-.00031	-.00075	-.00047	-.00076	-.00002	51	- 3	-122	-125	119
6.35	5.71	-.00027	-.00068	-.00048	-.00070	-.00005	54	- 8	-113	-121	105
7.62	5.71	-.00022	-.00060	-.00046	-.00063	-.00005	57	- 8	-101	-109	93
10.10	5.71	-.00019	-.00045	-.00040	-.00050	.00001	64	2	- 81	- 79	83
12.70	5.71	+.00008	-.00030	-.00033	-.00041	.00014	71	23	- 65	- 43	80
15.24	5.71	+.00030	-.00019	-.00031	-.00036	.00023	74	36	- 58	- 2	116

TABLE X.
PRINCIPAL STRESSES AND ASSOCIATED STRESS LOCI, DERIVED FROM MOHR'S CIRCLE OF STRAIN
CALCULATIONS USING X-RAY DATA

Δx	Δy	σ_x	ϵ_{45}	ϵ_y	ϵ_z	ϵ_1	σ_1	σ_2	$\sigma_1 - \sigma_2$	$\sigma_1 - \sigma_2$	$\sigma_1 - \sigma_2$
2.54	8.00	-.00040	-.00058	+.00005	-.00064	.00039	30	47	-103	-56	150
3.81	8.00	-.00035	-.00054	-.00013	-.00056	.00006	37	10	-90	-80	100
5.08	8.00	-.00031	-.00050	-.00024	-.00050	-.00003	41	-8	-81	-89	73
6.35	8.00	-.00026	-.00046	-.00028	-.00046	-.00008	46	-12	-74	-86	62
7.62	8.00	-.00022	-.00042	-.00026	-.00042	-.00006	48	-10	-55	-78	38
10.16	8.00	-.00017	-.00034	-.00020	-.00034	.00001	51	2	-73	-53	57
12.70	8.00	.00000	-.00026	-.00017	-.00028	.00011	58	18	-65	-27	63
15.24	8.00	+.00027	-.00019	-.00016	-.00027	.00038	60	61	-45	18	104
20.32	8.00	+.00040	-.00003	-.00015	-.00019	.00044	79	71	-31	69	102
25.40	8.00	+.00032	+.00004	-.00015	-.00015	.00032	85	52	-24	28	76
30.48	8.00	+.00016	-.00007	-.00015	-.00017	.00018	77	20	-27	2	56
2.54	9.53	-.00060	-.00051	+.00015	-.00070	.00025	19	40	-113	-73	153
3.81	9.53	-.00068	-.00041	+.00003	-.00069	.00006	7	10	-111	-101	121
5.08	9.53	-.00066	-.00033	+.00001	-.00049	.00004	14	6	-79	-73	85
6.35	9.53	-.00017	-.00038	-.00003	-.00033	.00013	36	21	-53	-32	74
7.62	9.53	-.00017	-.00032	-.00005	-.00033	.00011	37	18	-53	-35	71
10.16	9.53	-.00043	-.00029	-.00008	-.00043	-.00007	6	-11	-64	-80	58
12.70	9.53	-.00038	-.00026	-.00009	-.00048	-.00007	-4	-14	-77	-91	63
15.24	9.53	-.00031	-.00029	-.00010	-.00041	-.00010	6	-16	-50	-66	34

TABLE X.
PRINCIPAL STRESSES AND ASSOCIATED STRESS LOC1, DERIVED FROM MOHR'S CIRCLE OF STRAIN

Δx	Δy	e_x	e_{45}	e_y	e_z	σ_1	σ_2	σ_3	MN/m^2		
									σ_1	σ_2	$\sigma_1 - \sigma_2$
2.34	12.70	-.00038	-.00018	+.00020	-.00039	.00021	0	0	34	-63	-29
3.21	12.70	-.00032	-.00015	+.00020	-.00034	.00022	10	10	38	-35	-20
5.08	12.70	-.00056	-.00014	+.00020	-.00018	.00032	29	29	52	-29	-23
6.35	12.70	-.00007	-.00016	+.00020	-.00020	.00033	30	30	53	-32	-21
7.62	12.70	-.00010	-.00018	+.00020	-.00022	.00032	28	28	52	-35	-17
10.16	12.70	-.00002	-.00023	+.00020	-.00028	.00044	36	36	71	-40	-31
12.70	12.70	-.00060	-.00022	+.00017	-.00042	.00049	40	40	34	-68	-37
15.24	12.70	-.00010	-.00011	+.00012	-.00015	.00047	24	24	27	-24	3
2.34	22.86	-.00011	+.00010	+.00018	-.00012	.00019	-12	-12	31	-19	14
3.21	22.86	-.00009	+.00014	+.00027	-.00010	.00028	-8	-8	43	-16	29
5.08	22.86	-.00006	+.00016	+.00033	-.00006	.00035	-2	-2	56	-10	46
6.35	22.86	-.00004	+.00018	+.00043	-.00004	.00043	2	2	69	-6	63
7.62	22.86	-.00002	+.00020	+.00051	-.00002	.00051	5	5	82	-3	79
3.21	27.94	-.00018	+.00027	+.00028	-.00027	.00037	-22	-22	60	-43	17
5.08	27.94	-.00007	+.00022	+.00035	-.00008	.00036	-10	-10	58	-13	45
6.35	27.94	+.00001	+.00016	+.00043	0	.00044	8	8	71	0	71
7.62	27.94	+.00003	+.00011	+.00050	-.00002	.00053	17	17	89	-3	86

TABLE 8.
PRINCIPAL STRESSES AND ASSOCIATED STRESS LOGS, DERIVED FROM MOHR'S CIRCLE OF STRAIN
CALCULATIONS USING X-RAY DATA

Δx	Δy	ϵ_x	ϵ_{45}	ϵ_y	ϵ_1	ϵ_2	σ_x	σ_1	σ_2	$\sigma_1 + \sigma_2$	$\sigma_1 - \sigma_2$
1.27	31.75	-0.00012	+0.00031	+0.00020	-0.00027	+0.00035	-30	56	-43	13	99
2.34	31.75	-0.00014	+0.00025	+0.00026	-0.00022	+0.00034	-22	35	-35	26	90
3.81	31.75	-0.00015	+0.00018	+0.00032	-0.00017	+0.00024	-11	39	-27	12	66
5.08	31.75	-0.00016	+0.00012	+0.00038	-0.00016	+0.00038	-1	61	-26	35	87

TABLE XI.
PRINCIPAL STRESSES AND ASSOCIATED STRESS LOGS, DERIVED FROM MOHR'S CIRCLE OF STRAIN
CALCULATIONS USING X-RAY DATA

Δx	Δy	e_x	e_{45}	e_y	e_2	e_1	α_x°	σ_1	σ_2	$\sigma_1 + \sigma_2$	$\sigma_1 - \sigma_2$
								MN/m ²			
3.81	1.27	-.00036	-.00103	-.00101	-.00112	-.00047	66	- 72	-189	-252	108
3.81	2.34	-.00049	-.00101	-.00084	-.00103	-.00026	58	- 42	-169	-211	127
3.81	3.81	-.00043	-.00098	-.00067	-.00100	-.00013	53	- 16	-161	-177	165
3.81	5.08	-.00039	-.00092	-.00051	-.00092	.00002	49	3	-148	-145	151
3.81	6.35	-.00035	-.00084	-.00036	-.00084	.00013	45	21	-135	-114	156
3.81	7.62	-.00032	-.00075	-.00019	-.00075	.00024	41	39	-121	- 82	160
3.81	10.16	-.00027	-.00057	-.00012	-.00061	.00046	34	74	- 98	- 24	172
3.81	12.70	-.00022	-.00039	+.00019	-.00044	.00061	31	66	- 71	- 5	177
3.81	15.24	-.00018	-.00023	+.00026	-.00030	.00032	26	52	- 48	4	160
3.81	20.32	-.00013	+.00009	+.00023	-.00019	.00024	- 9	39	- 31	8	70
3.81	25.40	-.00018	+.00012	+.00026	-.00022	.00030	-16	48	- 35	13	83
3.81	30.48	-.00013	+.00005	+.00032	-.00018	.00032	2	52	- 29	23	81
2.34	1.27	-.00068	-.00120	-.00088	-.00121	-.00035	52	- 56	-195	-251	139
2.34	2.34	-.00060	-.00111	-.00066	-.00111	-.00015	47	- 24	-179	-203	155
2.34	3.81	-.00050	-.00102	-.00046	-.00102	.00006	44	10	-164	-154	174
2.34	5.08	-.00043	-.00093	-.00028	-.00093	.00022	41	33	-150	-115	185
2.34	6.35	-.00040	-.00084	-.00010	-.00086	.00034	38	55	-138	- 83	193
2.34	7.62	-.00040	-.00075	-.00002	-.00078	.00036	35	58	-126	- 68	184

TABLE XI,
PRINCIPAL STRESSES AND ASSOCIATED STRESS LOCL, DERIVED FROM MOHR'S CIRCLE OF STRAIN
CALCULATIONS USING X-RAY DATA

Δx	Δy	e_x	e_{45}	e_y	e_2	e_1	α_x^0	MN/m ²			$\sigma_1 + \sigma_2$	$\sigma_1 - \sigma_2$
								σ_1	σ_2	$\sigma_1 - \sigma_2$		
2.54	10.16	-.00038	-.00055	+.00017	-.00063	.00042	29	68	-101	-33	169	169
2.54	12.70	-.00034	-.00037	+.00018	-.00047	.00031	24	50	-76	-24	126	126
2.54	15.24	-.00030	-.00027	+.00019	-.00038	.00017	21	43	-61	-18	104	104
2.54	20.32	-.00018	+.00002	+.00020	-.00018	.00026	-1	32	-29	3	61	61
2.54	25.40	-.00022	+.00017	+.00021	-.00023	.00027	-20	43	-45	-2	85	85
2.54	30.48	-.00026	+.00002	+.00022	-.00025	.00024	-12	39	-45	-6	84	84
6.99	1.27	-.00013	-.00075	-.00068	-.00084	-.00001	64	-2	-135	-137	133	133
6.99	2.54	-.00017	-.00072	-.00064	-.00080	-.00001	63	-2	-129	-131	127	127
6.99	5.08	-.00017	-.00061	-.00050	-.00066	-.00003	60	-2	-106	-108	104	104
6.99	7.62	-.00017	-.00050	-.00031	-.00051	.00003	53	5	-82	-77	87	87
6.99	10.16	-.00014	-.00038	-.00002	-.00039	.00023	39	37	-63	-26	100	100
6.99	15.24	-.00005	-.00016	+.00006	-.00022	.00053	28	85	-35	50	120	120
6.99	20.32	-.00004	0	+.00048	-.00012	.00056	20	90	-19	71	109	109
6.99	25.40	-.00001	+.00004	+.00050	-.00008	.00057	19	92	-13	79	105	105
6.99	30.48	-.00013	-.00004	+.00047	-.00020	.00054	17	89	-32	57	121	121

TABLE XI.
PRINCIPAL STRESSES AND ASSOCIATED STRESS LOU, DERIVED FROM MOHR'S CIRCLE OF STRAIN
CALCULATIONS USING X-RAY DATA

Δx	Δy	MN/m ²									
		e_x	e_{45}	e_y	e_2	e_1	α_0	σ_1	σ_2	$\sigma_1 + \sigma_2$	$\sigma_1 - \sigma_2$
22.86	1.27	+0.0041	+0.0009	-0.0038	-0.00079	.00042	85	68	-63	5	131
22.86	2.54	+0.0032	+0.0011	-0.0033	-0.0013	.00034	80	55	-56	-	111
22.86	5.08	+0.0028	+0.0016	-0.0022	-0.0025	.00031	76	50	-40	10	90
22.86	7.62	+0.0026	+0.0020	-0.0015	-0.0020	.00031	72	50	-32	-8	82
26.16	5.08	+0.0025	+0.0022	-0.0028	-0.0041	.00038	114	61	-66	-	127
26.16	7.62	+0.0030	+0.0017	-0.0021	-0.0024	.00033	77	53	-39	14	92
31.75	1.27	+0.0030	+0.0029	-0.0038	-0.0051	.00043	68	69	-82	-13	151
31.75	2.54	+0.0026	+0.0024	-0.0030	-0.0040	.00036	69	58	-64	-	122
31.75	5.08	+0.0019	+0.0011	-0.0017	-0.0020	.00022	75	35	-32	3	67

FIGURE 1

DIAGRAM OF A DZIL BIT

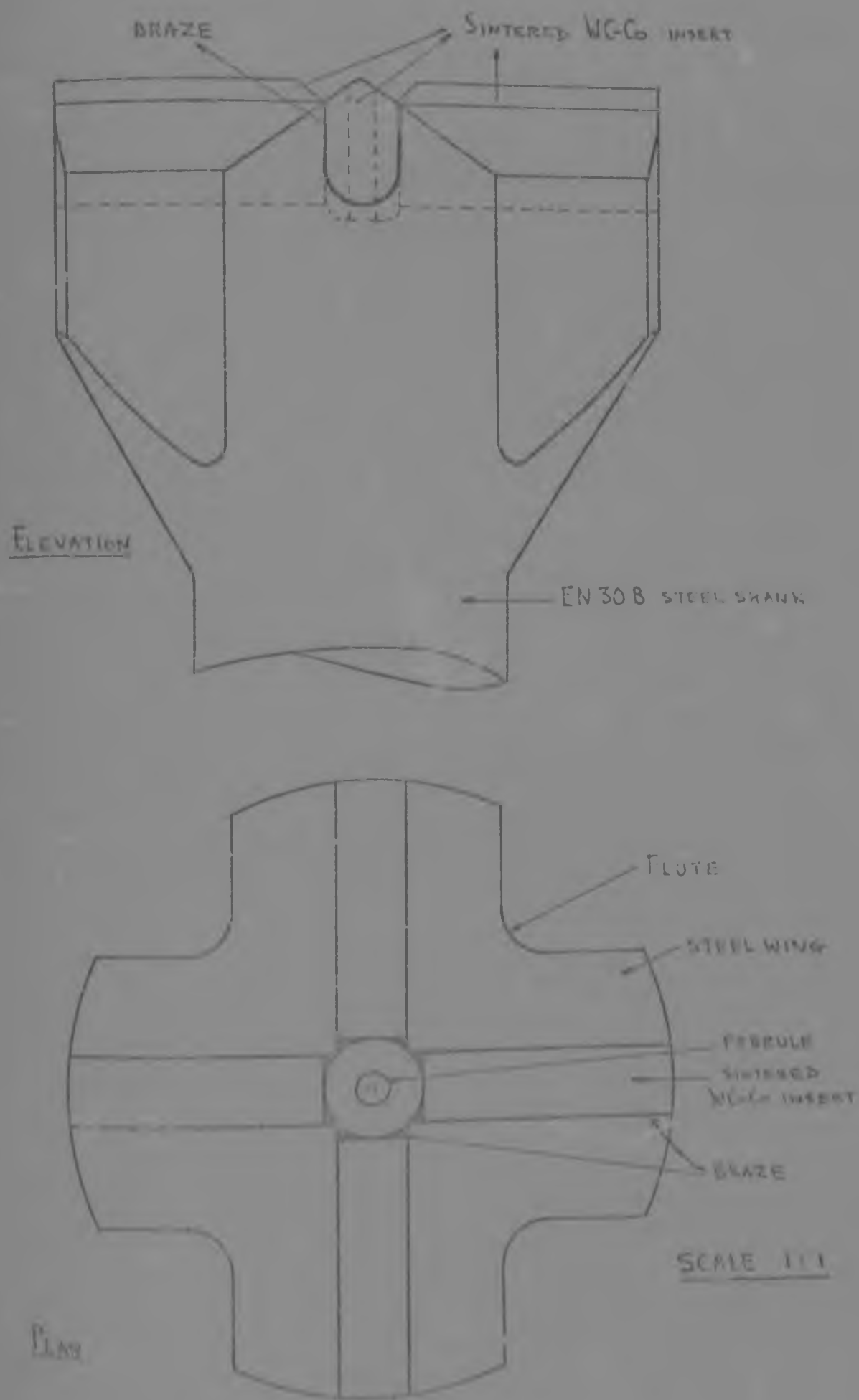


Figure 4. THE EFFECT OF SECTIONING AND METALLOGRAPHIC PREPARATION ON THE (310) DIFFRACTION PEAK OF PEARLITE

NOTE: For fairer comparison and to avoid the need to superimpose various peaks, the vertical scales for relative intensities have been moved downward by 10 per cent for each successive curve. The graph plotted for Reference Annealed Pearlite is on the same scale as that for the electro-polished sample, but the electro-polished specimen peak has been displaced to the right hand side.

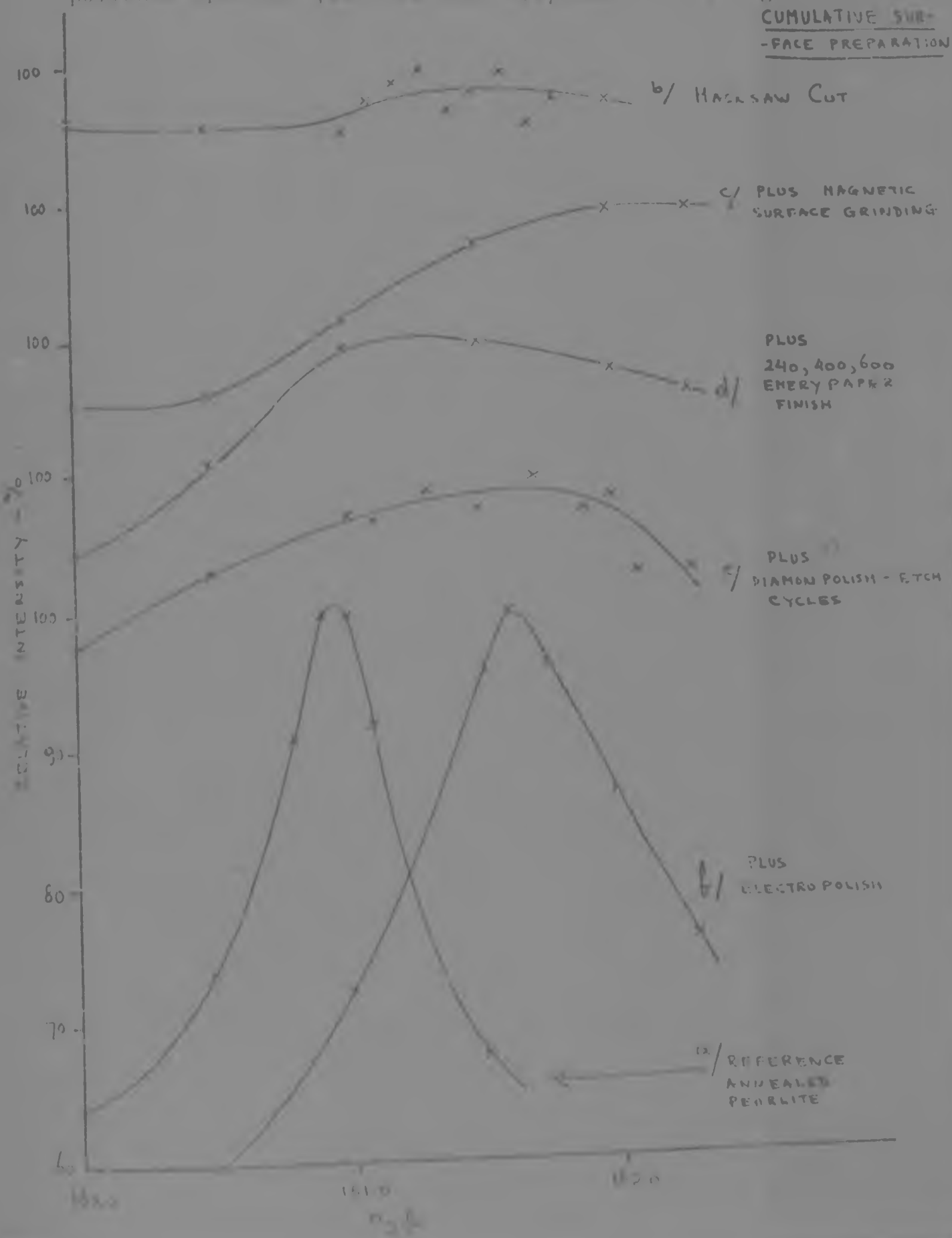


FIGURE 5. THE POLARIZATION OF AN ELECTROLYTICALLY STRENGTHENED SPECIMEN TO WHICH A 1000 STRAIN-GAUGE HAS BEEN ADDED



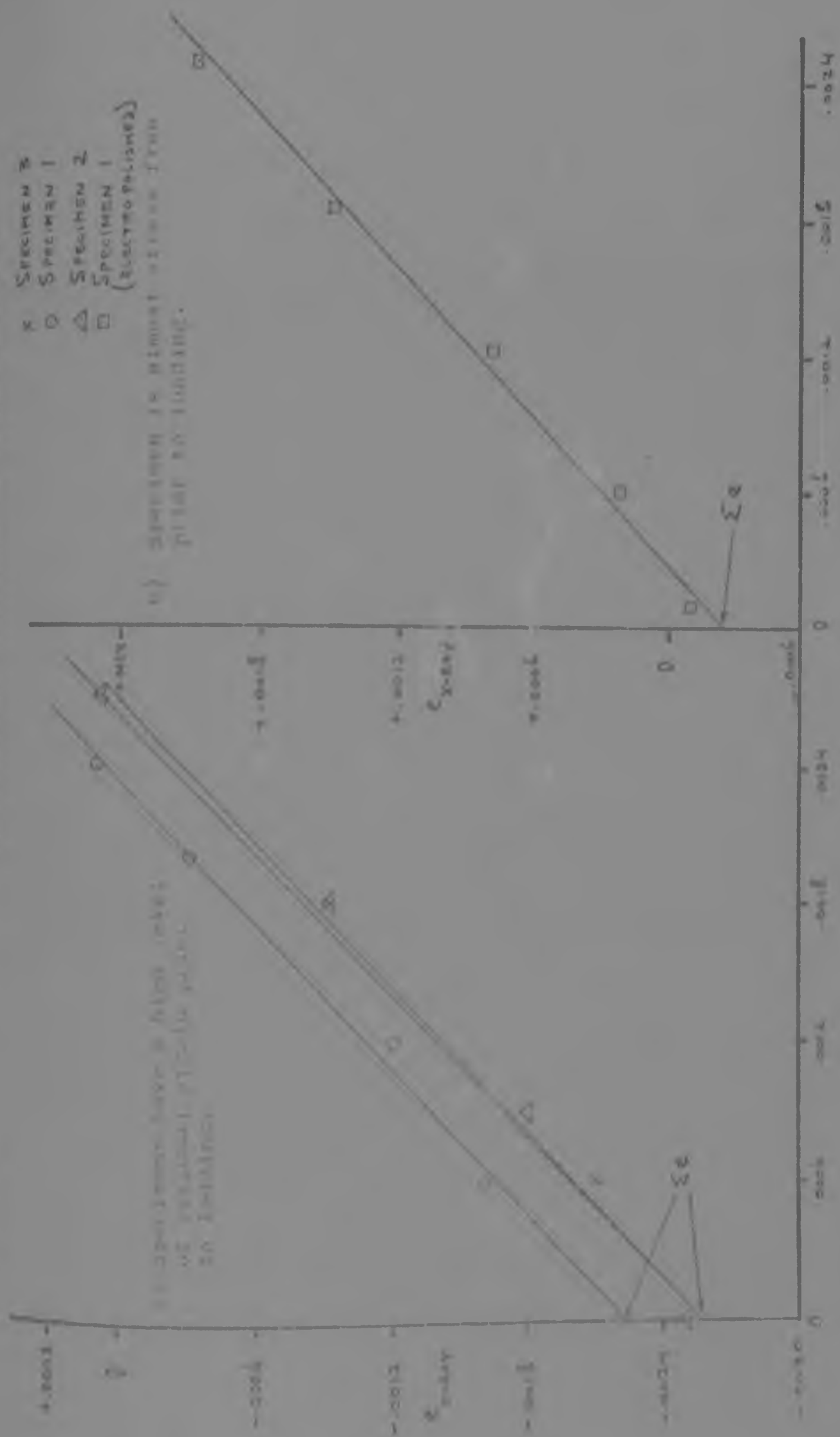
A AND B ARE FIXED JOINTS AT B AND C 2 STEEL PLUNGERS TRANSMIT FORCE TO SPECIMEN



FIGURE 6. PLAN

Figure 5

the maximum average strain is indicated by the top of the curve in Figure 5. The maximum average strain is indicated by the top of the curve in Figure 5.



Extremely Average Strain (4 Point Bending)

Figure 7

Plot of $\sin^2 \psi$ vs d for various values of λ and θ . The curves are calculated for $\lambda = 0.1$ nm and $\theta = 0.1$ rad.

Plot of $\sin^2 \psi$ vs d for various values of λ and θ . The curves are calculated for $\lambda = 0.1$ nm and $\theta = 0.1$ rad.

Legend for Symbols Applied in Figure 7:

- $\lambda = 0$
- $\lambda = 0.0001$
- $\lambda = 0.001$
- $\lambda = 0.002$
- $\lambda = 0.004$

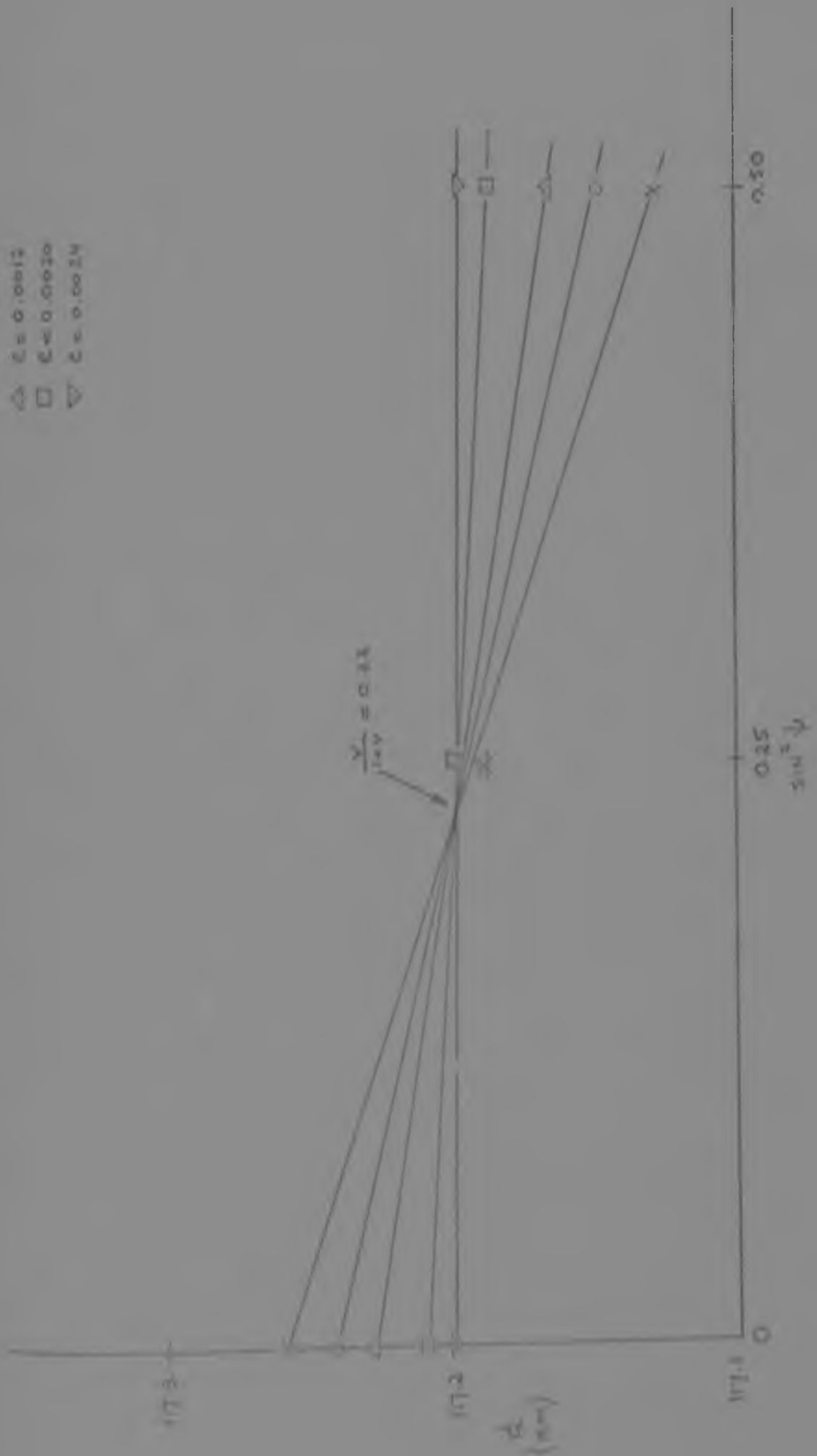


Figure 8

Comparison of the (a) and (b) curves shows the difference in the effect of the (a) and (b) curves on the (c) and (d) curves.

1. Unburned Fuel Gas
2. Sintered Compact
3. Sintered and Tempered Compact
4. Sintered and Cold Blasted Compact
5. Sintered and Electrically Ground Compact
6. Sintered and Ground Compact
7. Sintered and Vapour Blasted Compact



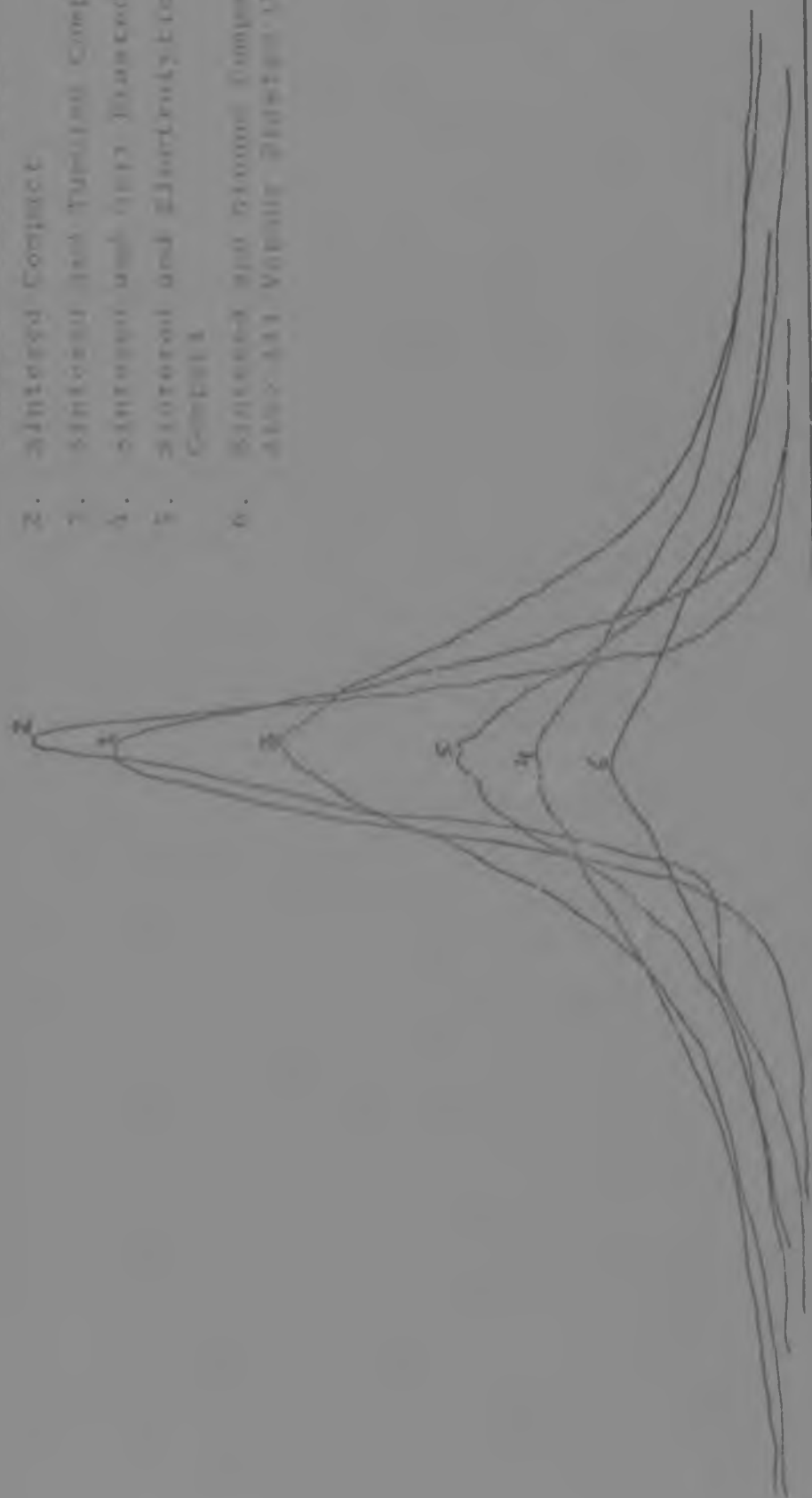
$\Delta t = 0.5^\circ$

DOUBLE DIFFRACTION ANGLE 2θ

FIGURE 9

CONTOURS OF THE 11001 DIFFRACTION PEAK OF TUNGSTEN CARBIDE IN COMPACTS AFTER SEVERAL DIFFERENT SURFACE PREPARATION TECHNIQUES

1. Tungsten Carbide Powder
2. Sintered Compact
3. Sintered and Tumbled Compact
4. Sintered and 4000 Grit Bounced Compact
5. Sintered and Electrolytically Ground Compact
6. Sintered and Ground Compact also 4000 Grit Bounced Compact



$$\Delta 2\theta = 0.5^\circ$$

DIFFRACTION ANGLE 2θ

Figure 10.

CONTOUR OF THE 1000 DEFLECTION VOLT OF TUNGSTEN CATHODE IN CONTACTS AFTER
SEVERAL HYPERTHERMAL DEGRADATION TREATMENTS

1. Tungsten Cathode (new)
2. Sintered Cathode
3. Sintered and Tungsten Cathode
4. Sintered and Tungsten Cathode
5. Sintered and Tungsten Cathode
6. Sintered and Tungsten Cathode



0.25 x 0.15

Double Deflection Angle 2.8

FIGURE II.

POSITIONS OF THE 100% DIFFRACTION PEAK OF TUNGSTEN CARBIDE IN SINTERED
COMPACTS AND FRESH POWDER

1. Tungsten Carbide Powder
2. Sintered Compact
3. Sintered and Tumbled Compact
4. Sintered and prepared as 4, 5 and 6
in Figure 10, 11 and 12.



$2\theta 40.5^\circ$
DOUBLE DIFFRACTION ANGLE 2θ

FIGURE 12
 POSITIONS OF THE (112) DIFFRACTION PEAK OF TUNGSTEN CARBIDE IN COMPACTS AND
 POWDER AT ROOM TEMPERATURE

1. Tungsten Carbide powder
 2. Sintered Compact
 3. Sintered and Tumbled Compact
- Numbers 4, 5 and 6 showed nearly any
 peak shift.



$\Delta 2\theta = 0.5^\circ$
 DOUBLE DIFFRACTION ANGLE 2θ

FIGURE 13

DETERMINATION OF PARTICLE SIZE AND PLASTIC STRAIN FROM
DIFFRACTION PEAK BROADENING

REFERENCE SPECIMEN: ANNEALED MC-BORON

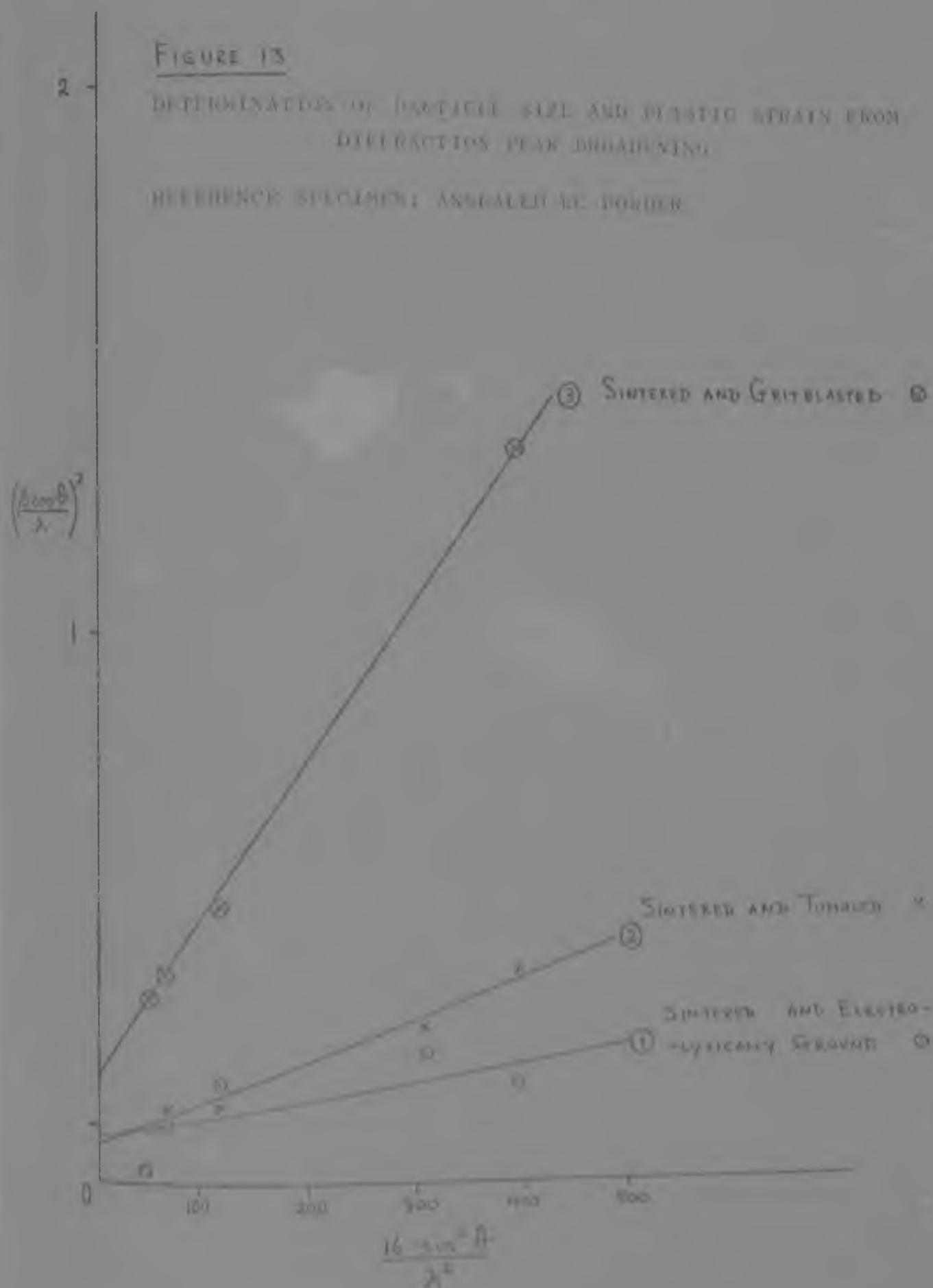
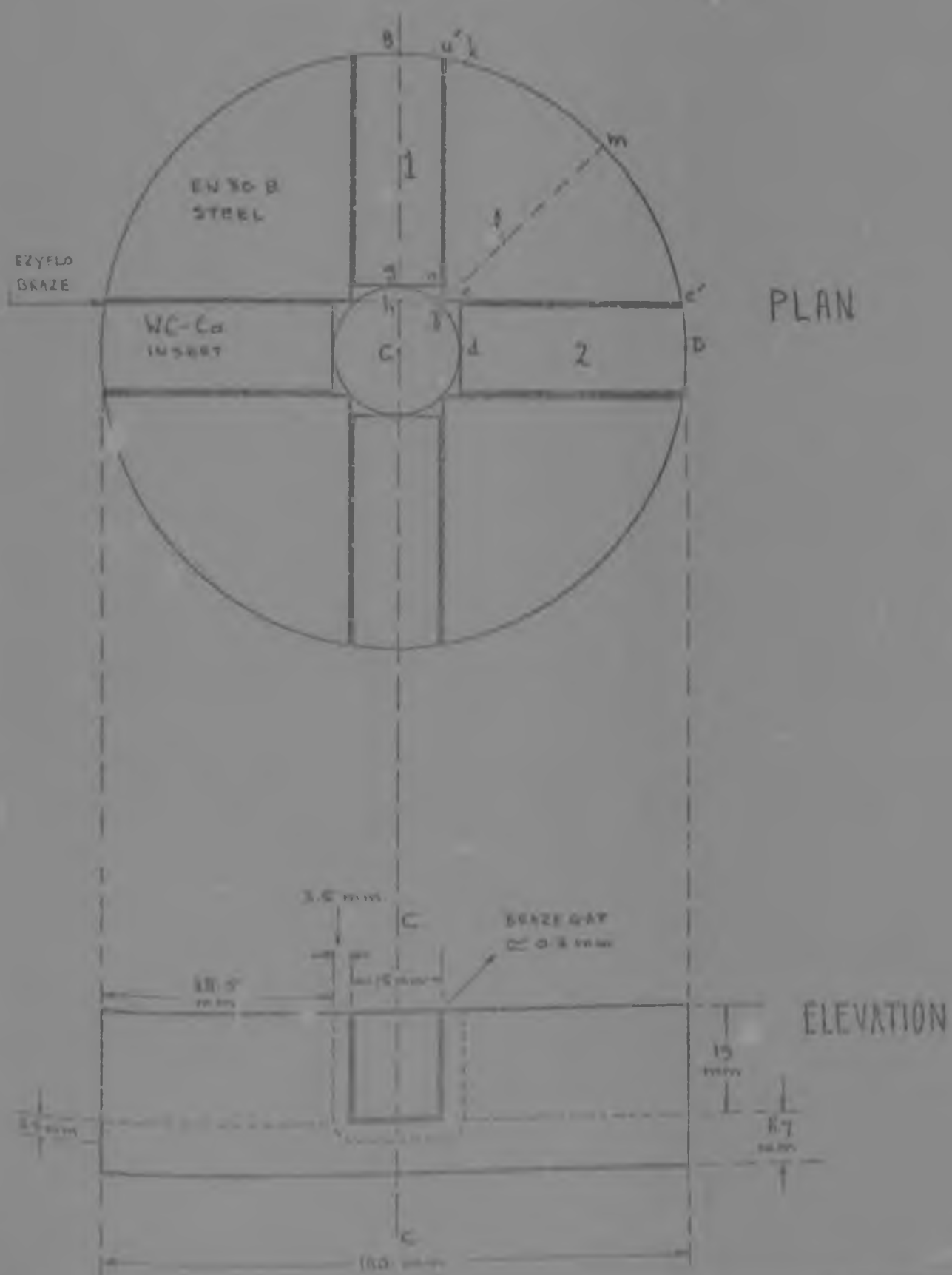


FIGURE 14.

D/D HIT ANALOGUE, PLAN AND ELEVATION.

SCALE: 1 : 1



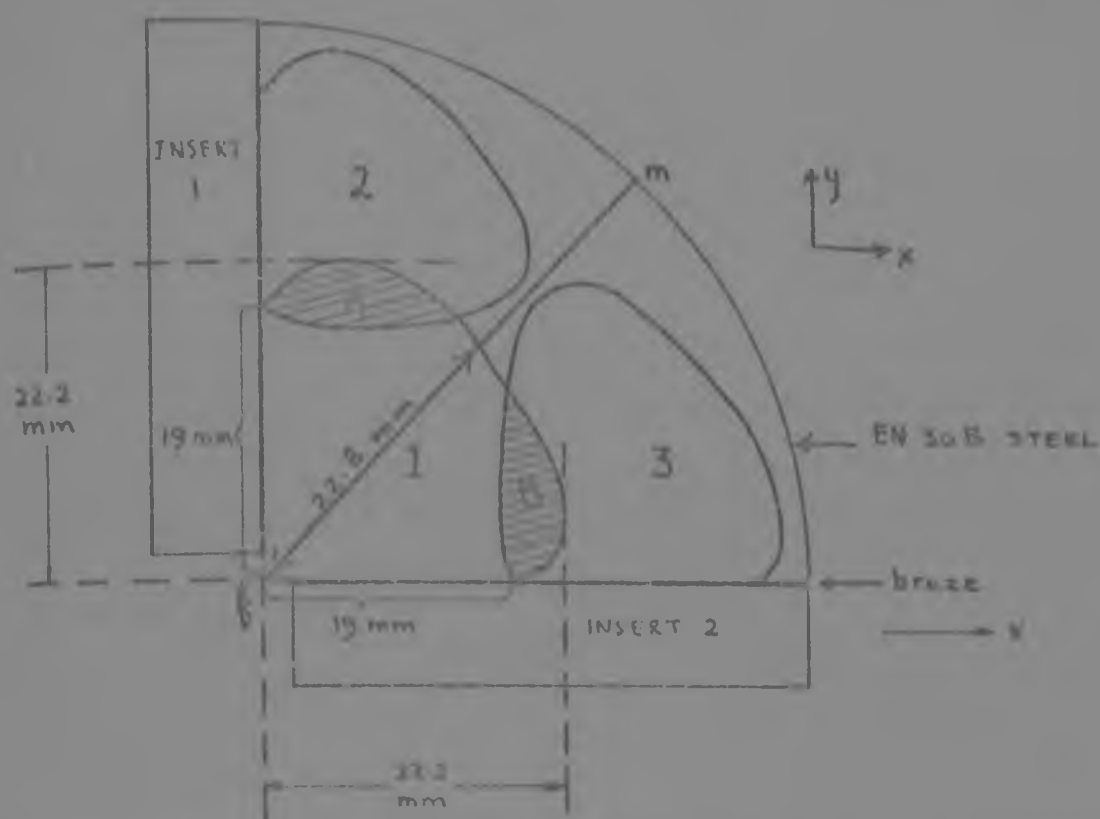


FIGURE 15. ELECTRODEPOSITED AREAS ON SECTOR SURFACES. THE THREE ELECTRODEPOSITED AREAS ARE DESIGNATED 1, 2 AND 3.

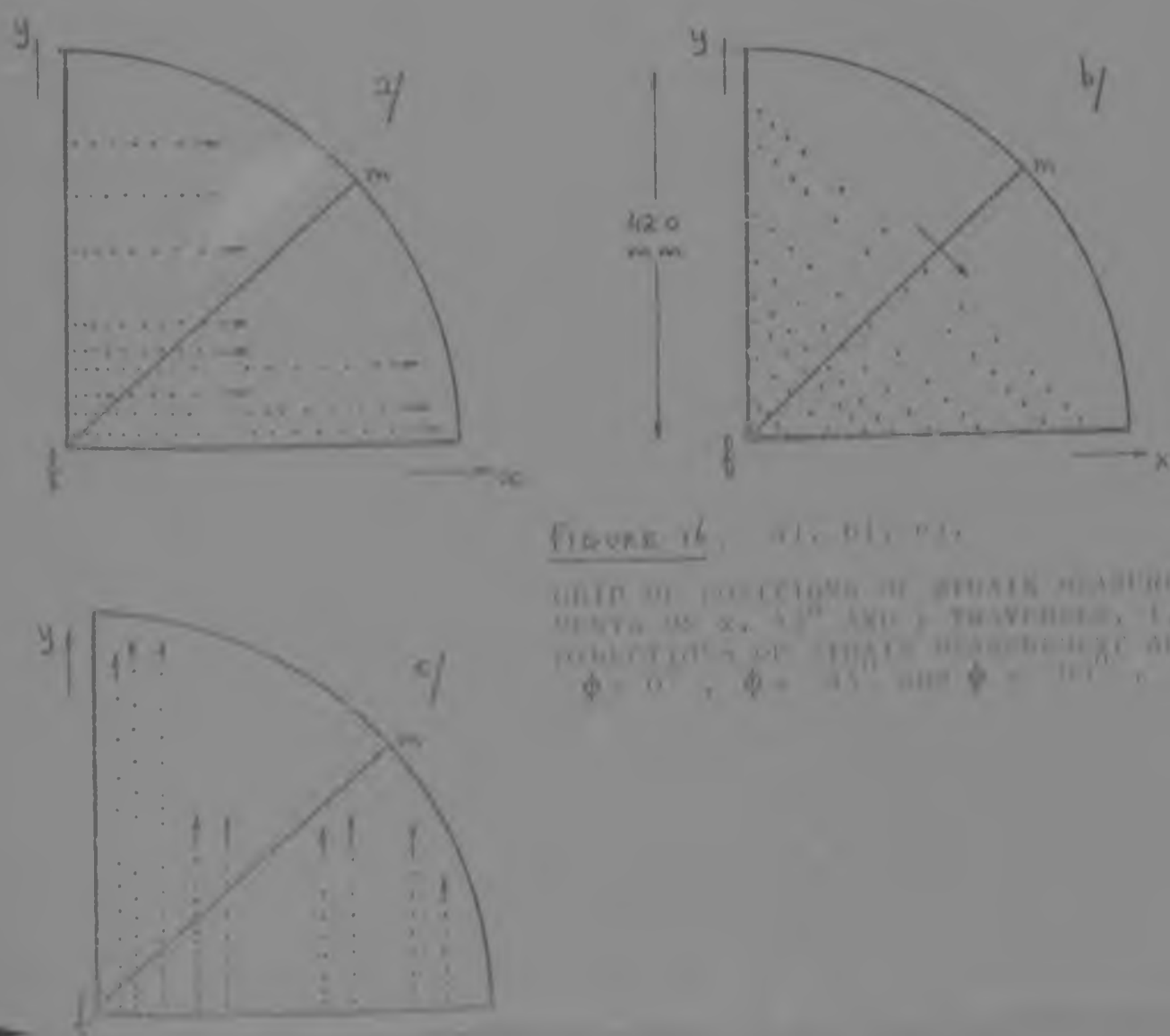


FIGURE 16. STRAIN MEASUREMENTS.

GRID OF LOCATIONS OF STRAIN MEASUREMENTS ON x , y AND z TRAVERSES, (a) LOCATIONS OF STRAIN MEASUREMENTS ON x AND y TRAVERSES, (b) LOCATIONS OF STRAIN MEASUREMENTS ON x AND y TRAVERSES, (c) LOCATIONS OF STRAIN MEASUREMENTS ON x AND y TRAVERSES.

Figure 17

STRAIN ϵ FOR σ - ORIENTATION MEASURED ALONG THE TRAVERSE $\Delta y = 1.27$ mm

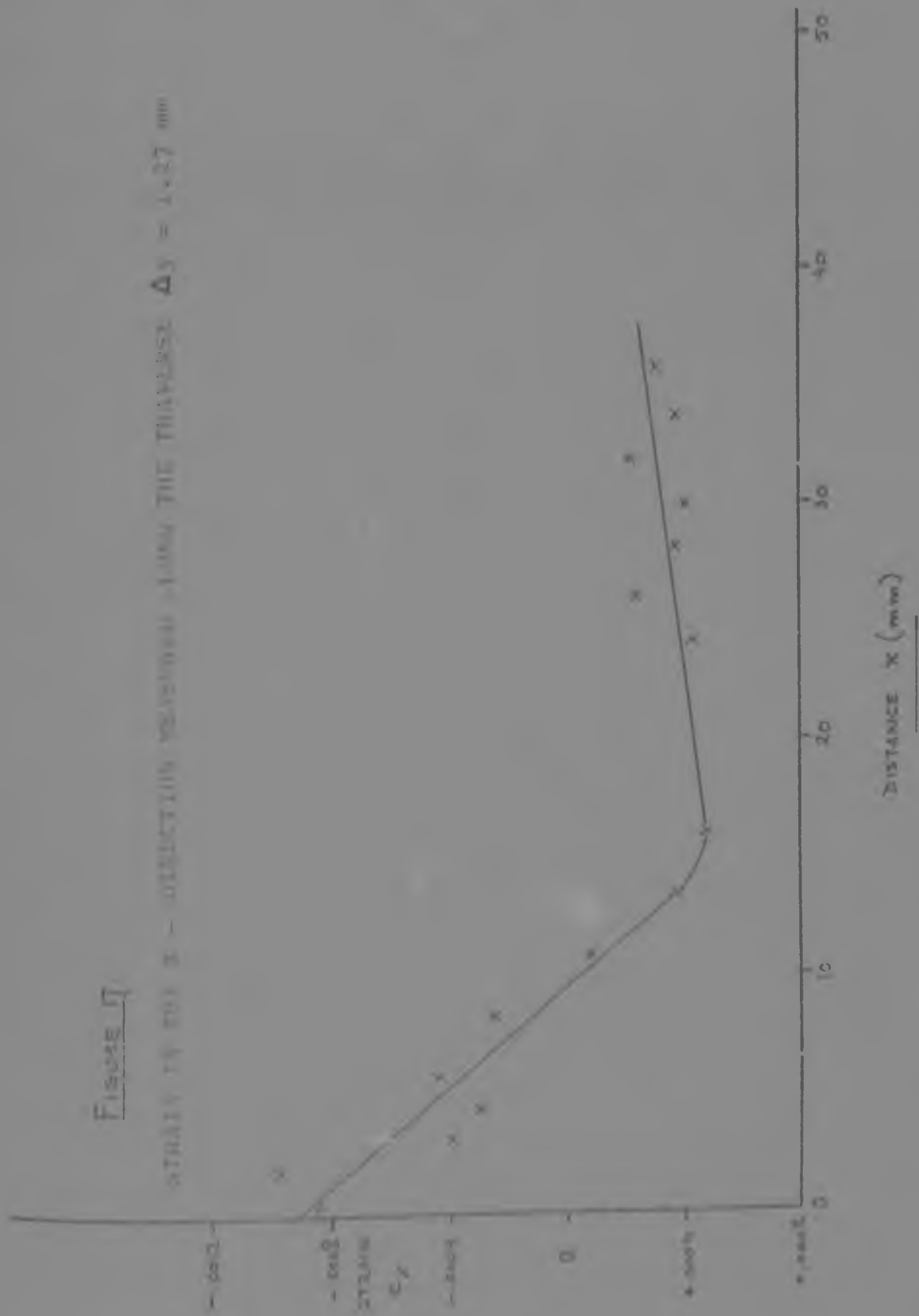


Figure 18.

STRAIN IN THE X-DIRECTION MEASURED ALONG THE TRAVERSE $\Delta_1 = 3.81 \text{ mm}$

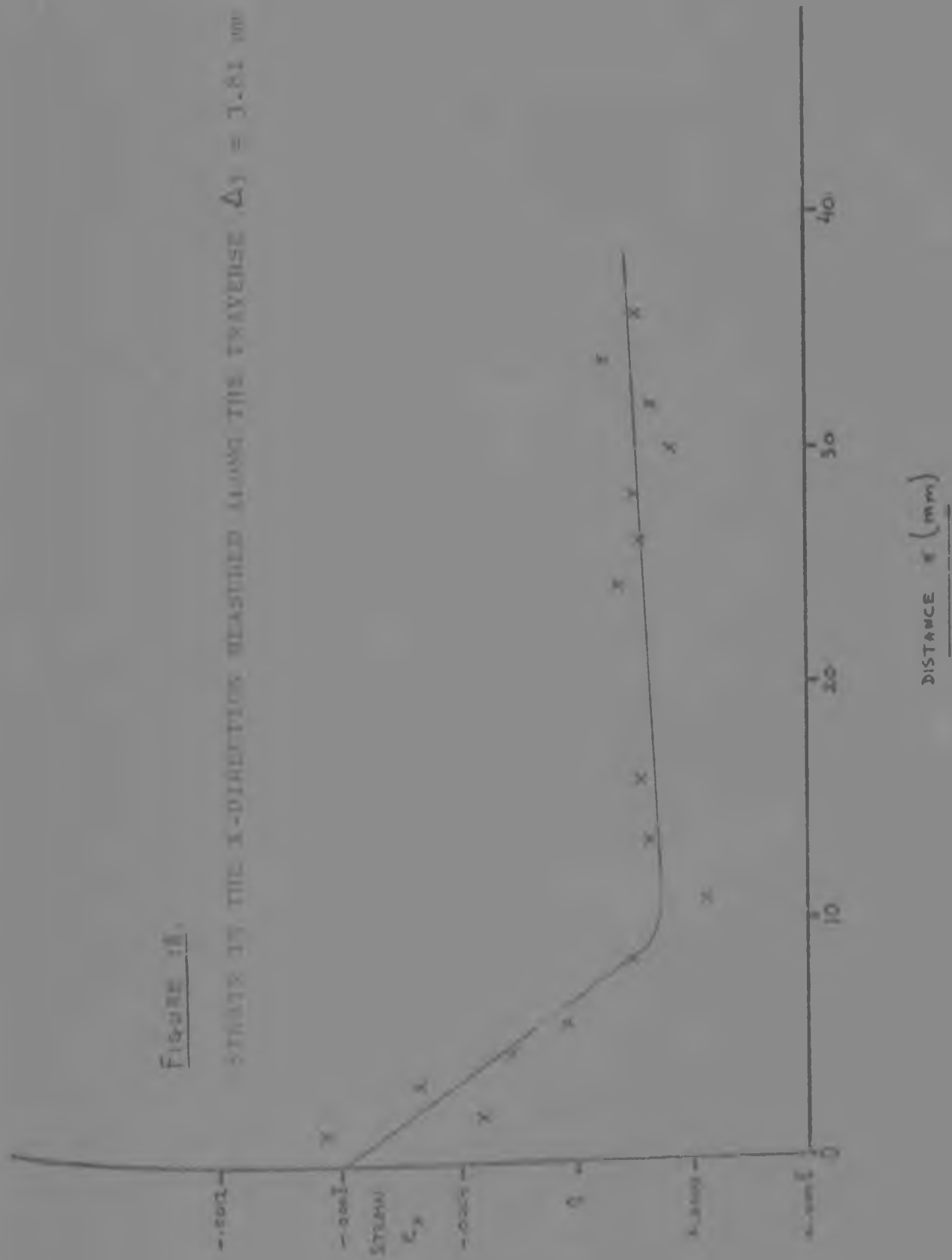


Figure 13.

100% IS THE X-DIRECTION MEASURED ALONG THE TRANSVERSE $\Delta y = 5.71$ mm

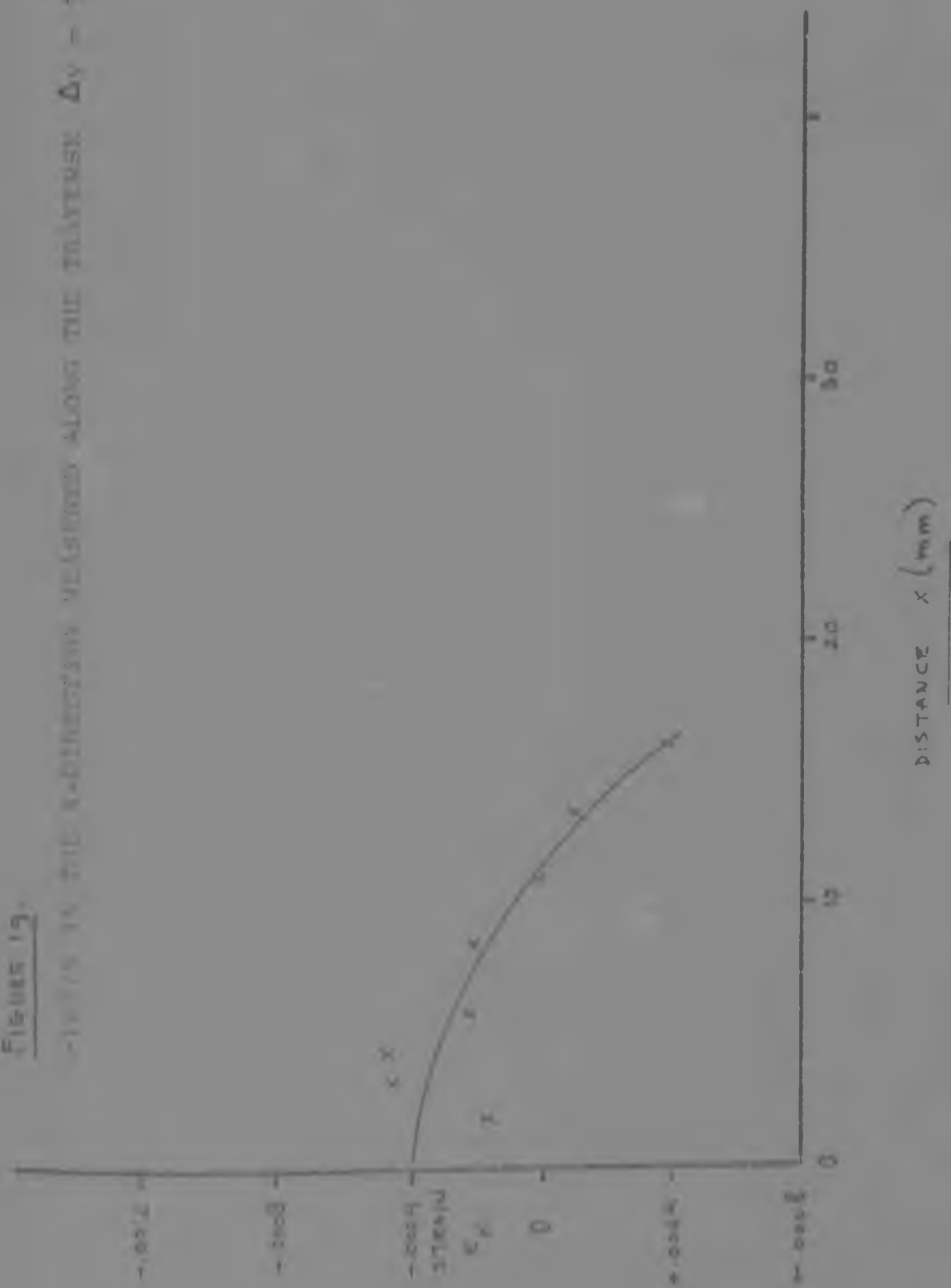
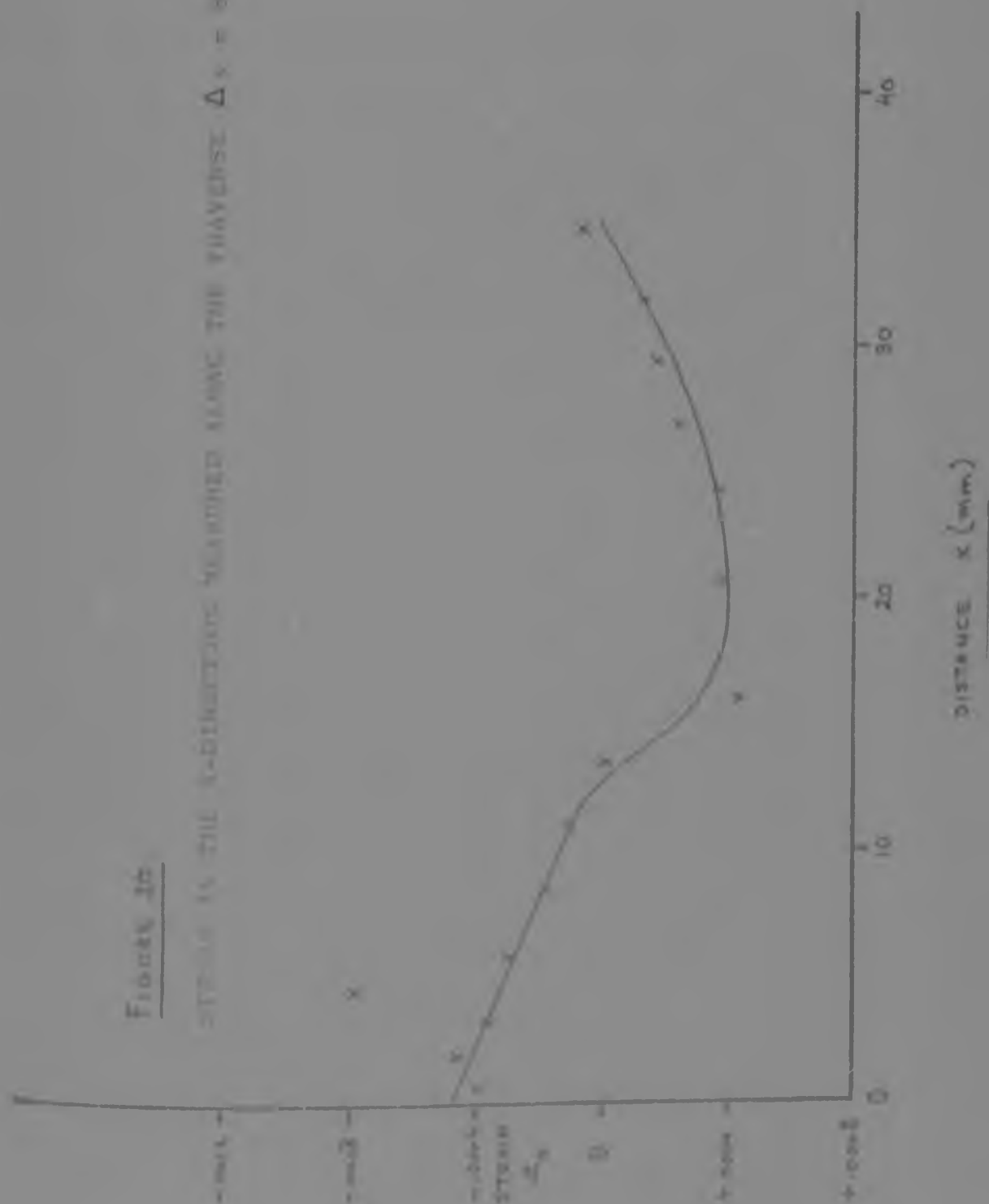


Figure 3a.

STATION IS THE RADIIATION MEASURED ALONG THE TRAVERSE $\Delta x = 8.00$ cm



STRAIN IN THE X-DIRECTION MEASURED ALONG Δy TRAVERSES

Figure 21

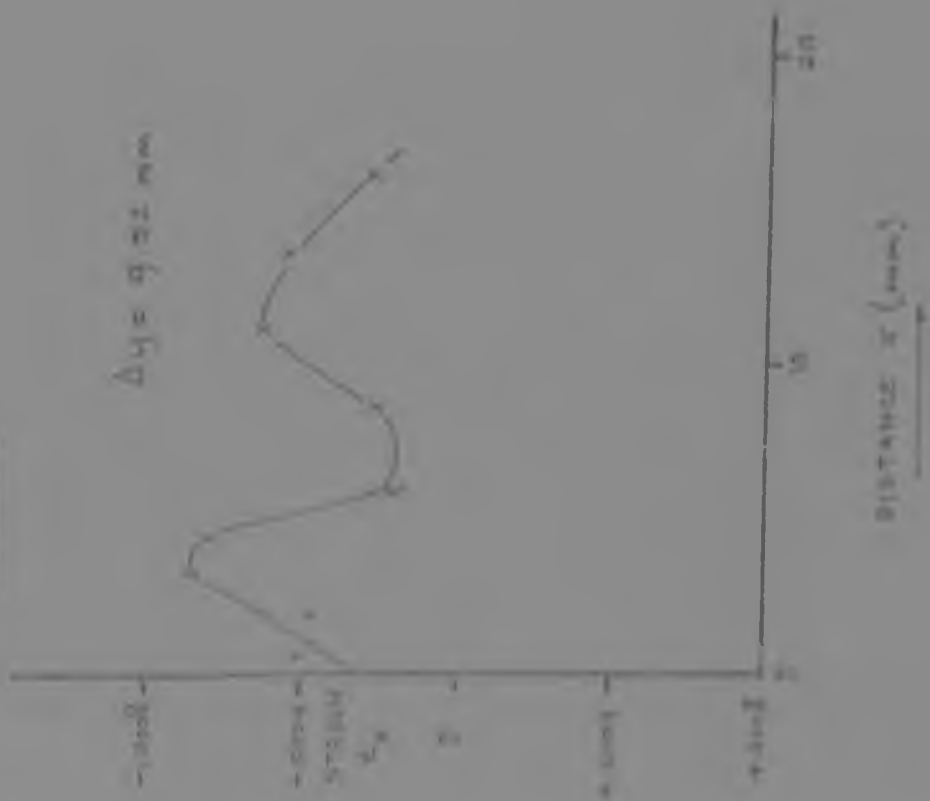


Figure 22

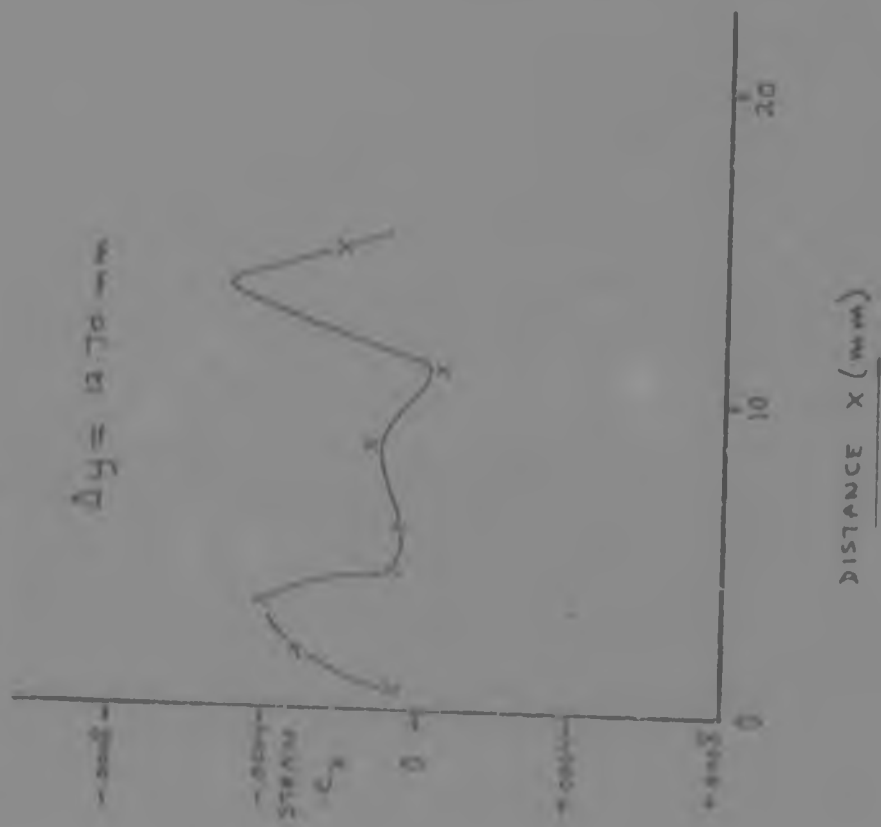


Figure 23. The x-axis represents distance along the trajectory $\Delta y = 22.84 \text{ mm}$ $\Delta y = 27.93 \text{ mm}$

Figure 23.

$$\Delta y = 22.84 \text{ mm}$$

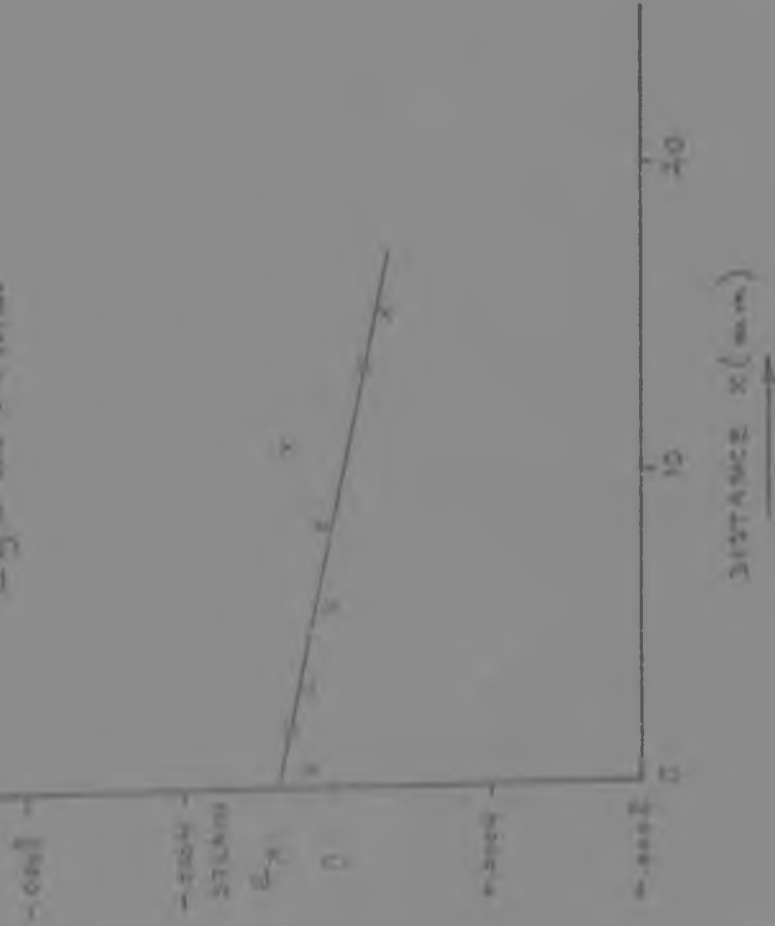


Figure 24.

$$\Delta y = 27.93 \text{ mm}$$

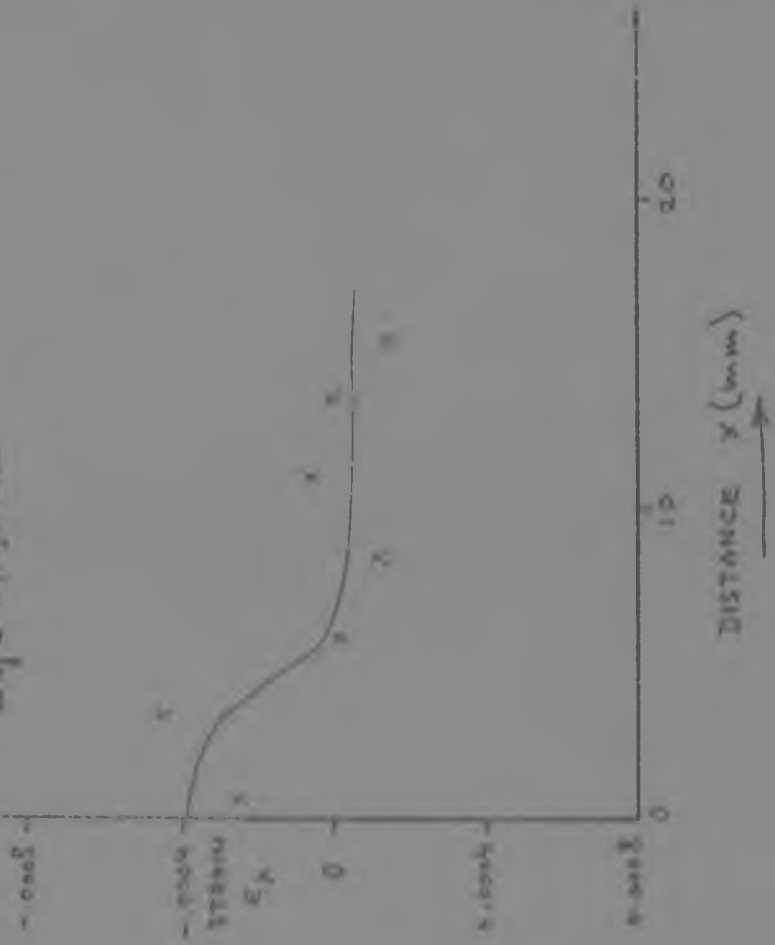
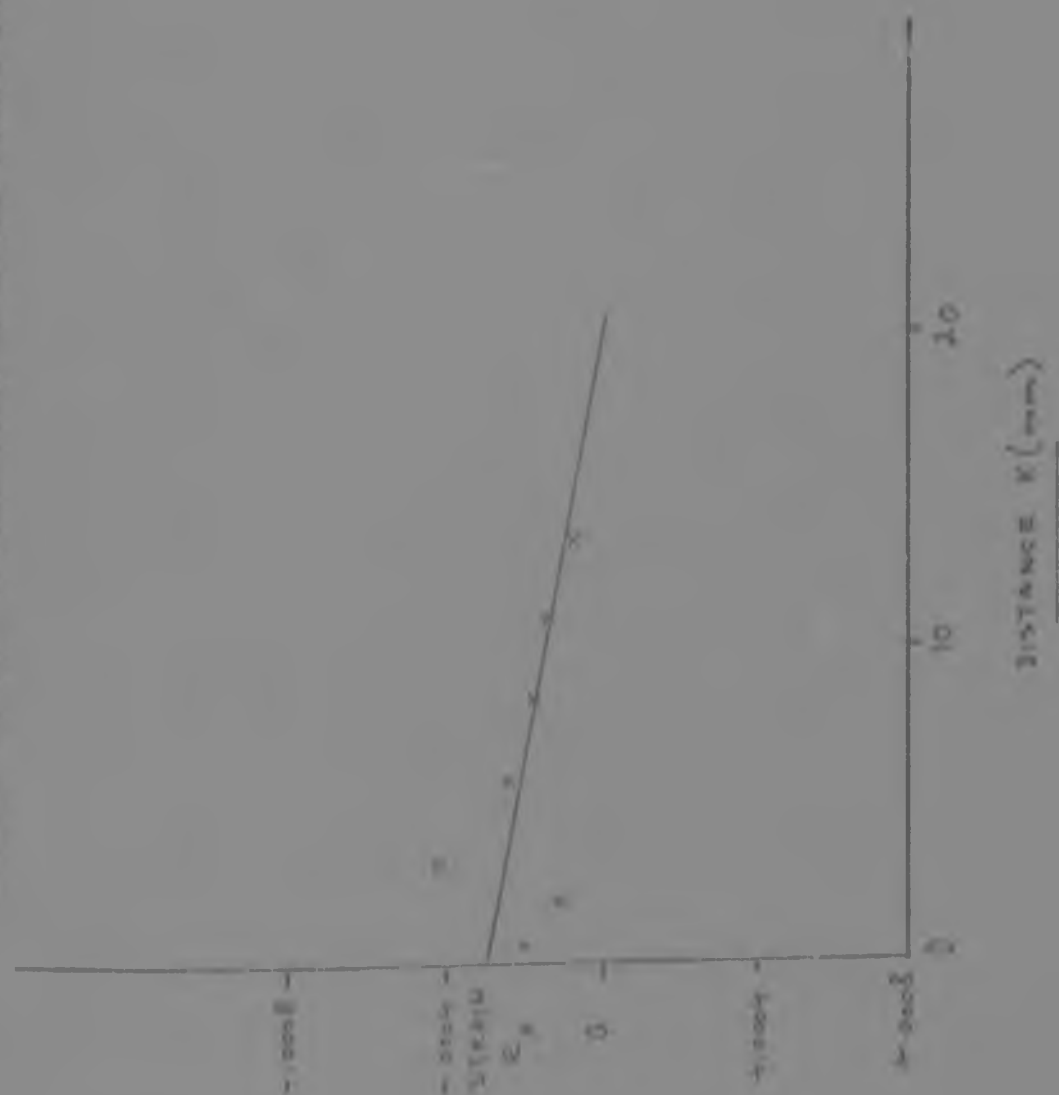


Figure 25.

SCOTS IN THE X-DIRECTION MEASURED ALONG THE TRAVERSE $\Delta y = 31.78 \text{ mm}$



The following data were obtained from the tests shown in Figures 26, 27, and 28. The distance b is the distance from the point of application of the load to the support.

Figure 26.

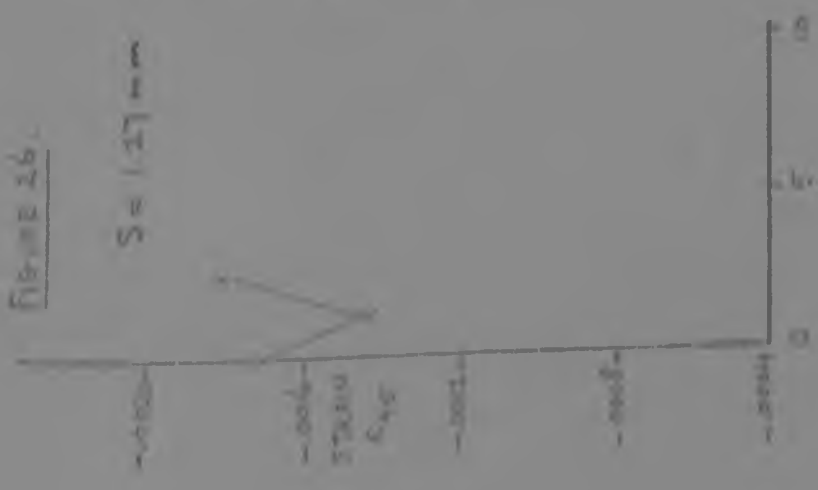


Figure 27.

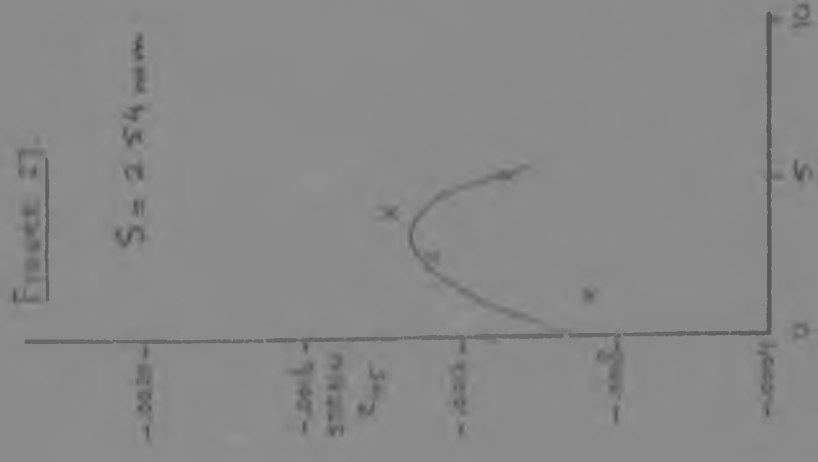
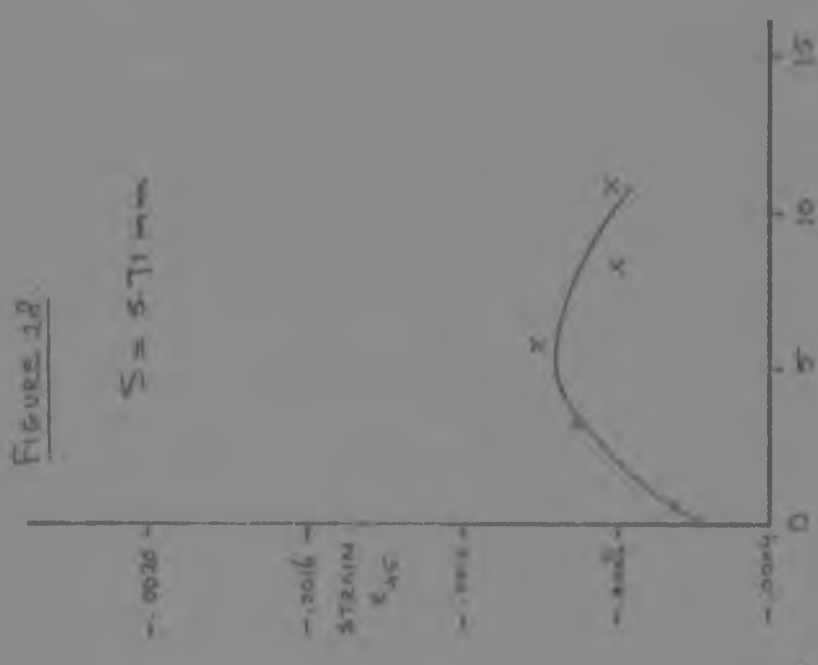


Figure 28.



DISTANCE b (mm) $\rightarrow$

A 10.00 mm thick plate of 2024-T3 aluminum alloy is tested in a tensile testing machine. The load is applied at a constant rate of 0.25 mm/min. The load is applied at a constant rate of 0.25 mm/min. The load is applied at a constant rate of 0.25 mm/min.

Figure 29

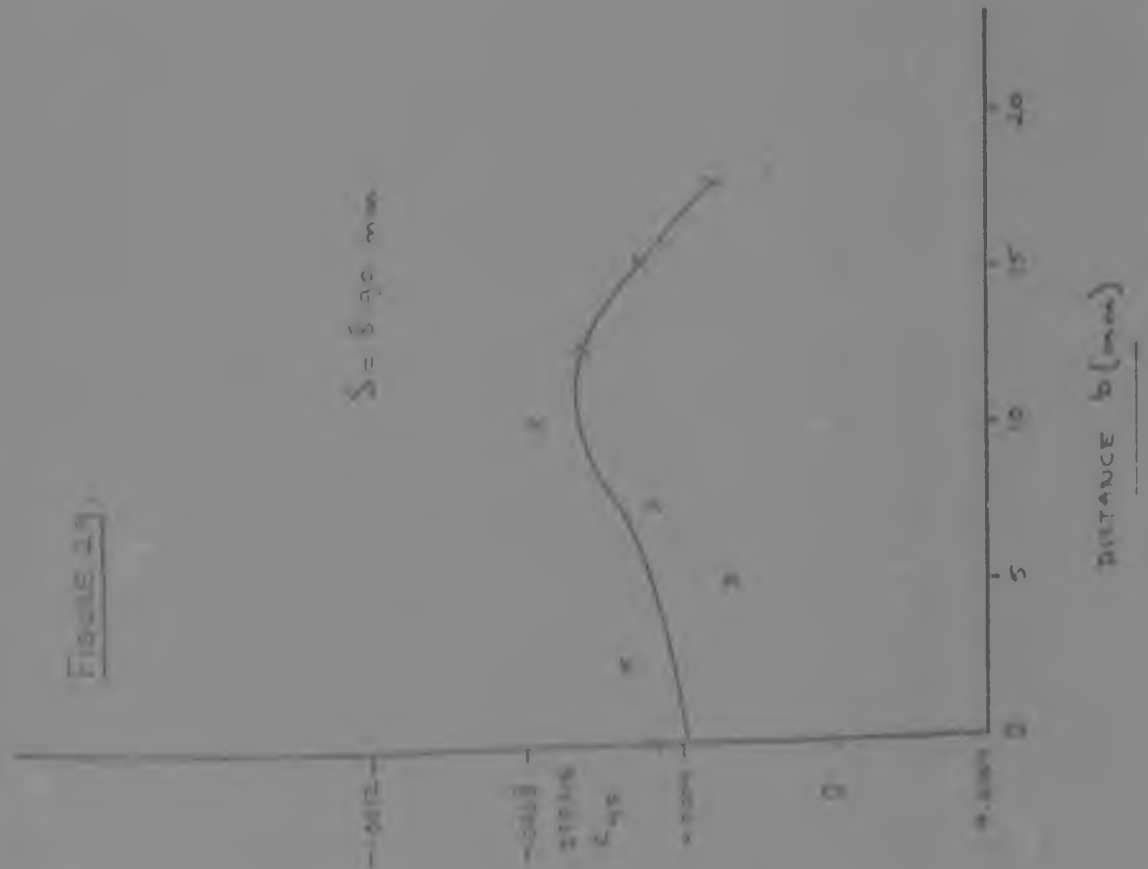


Figure 30

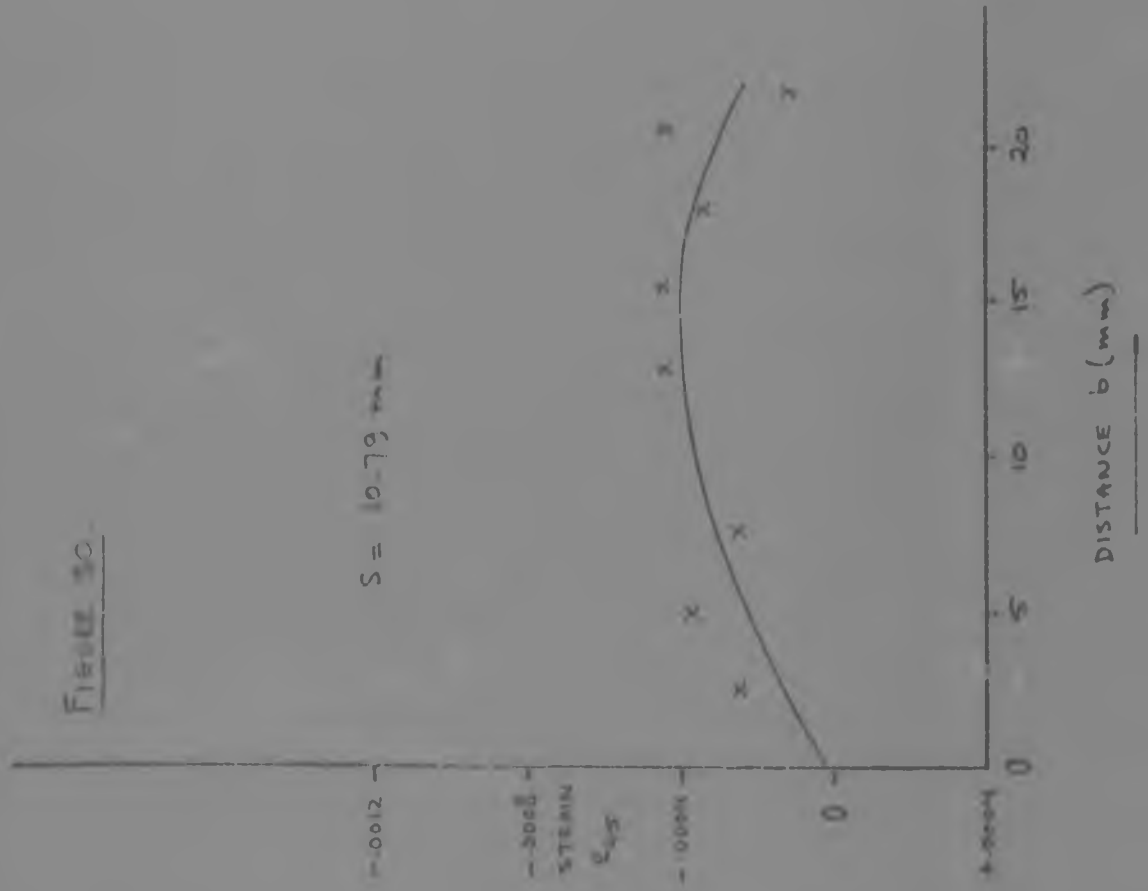


Figure 31

GRAPH OF THE $\log$ CONCENTRATION PLASMA ALONG THE TRAJECTORY
AT $S = 6.0-20$ mm FROM POINT X

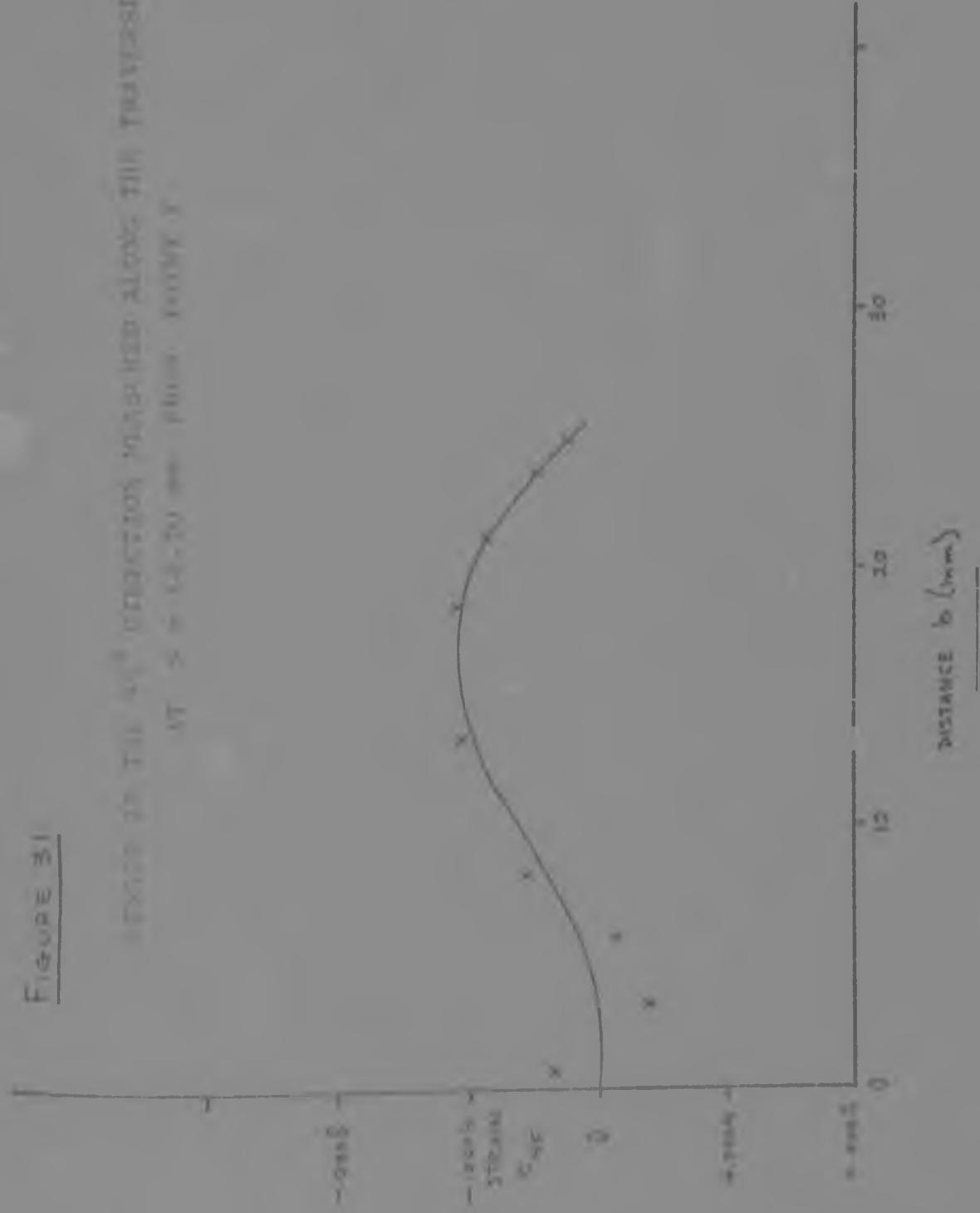


Figure 32

SWAY IN THE 60° DIRECTION MEASURED ALONG THE TRAVERSE
 AT $S = 17.78$ mm FROM POINT 1

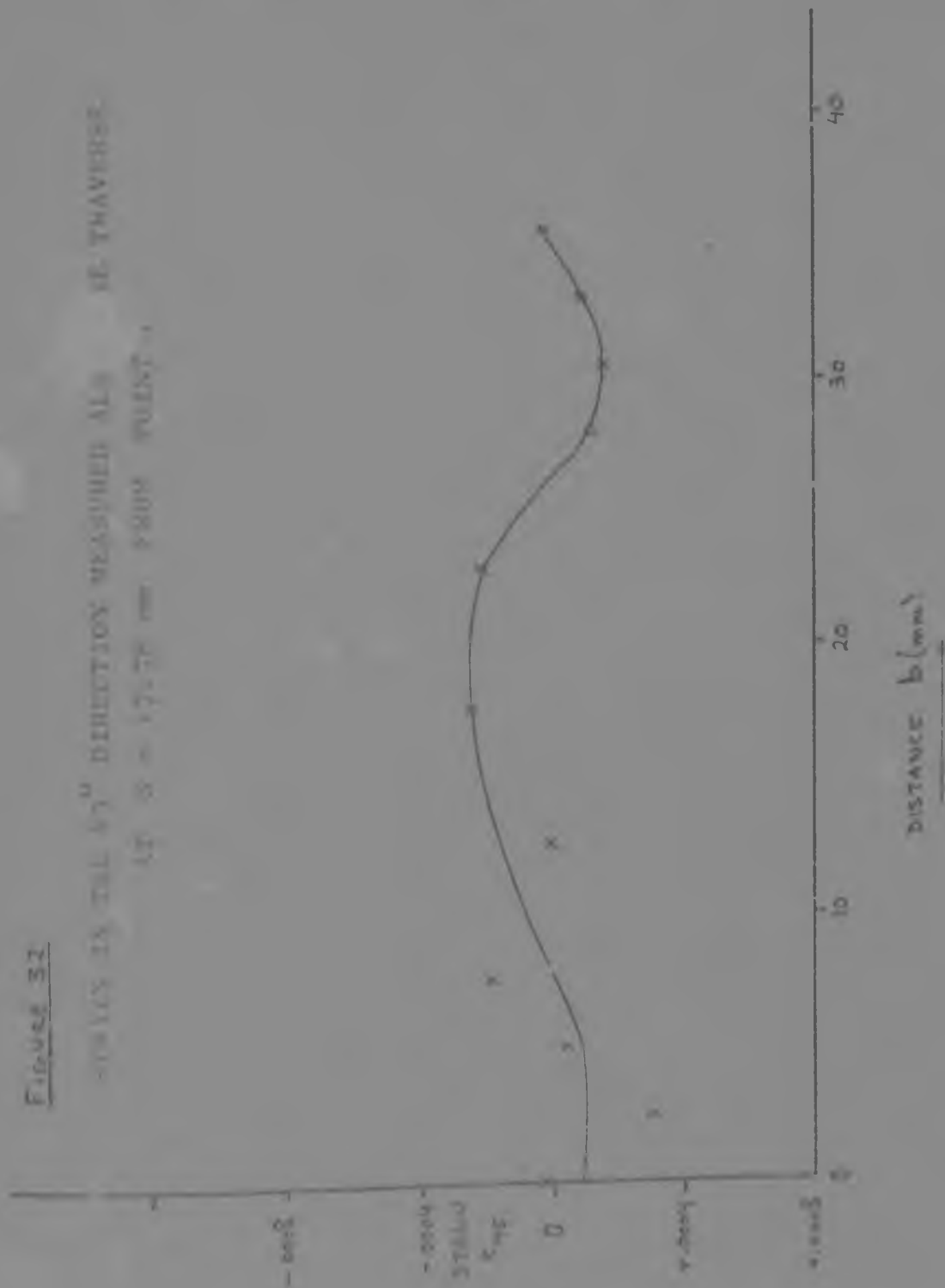


Figure 25

STRESS IN THE 45° DIRECTION MEASURED ALONG THE TRANSVERSE
 AT $b = 0.25$ mm FROM POINT P

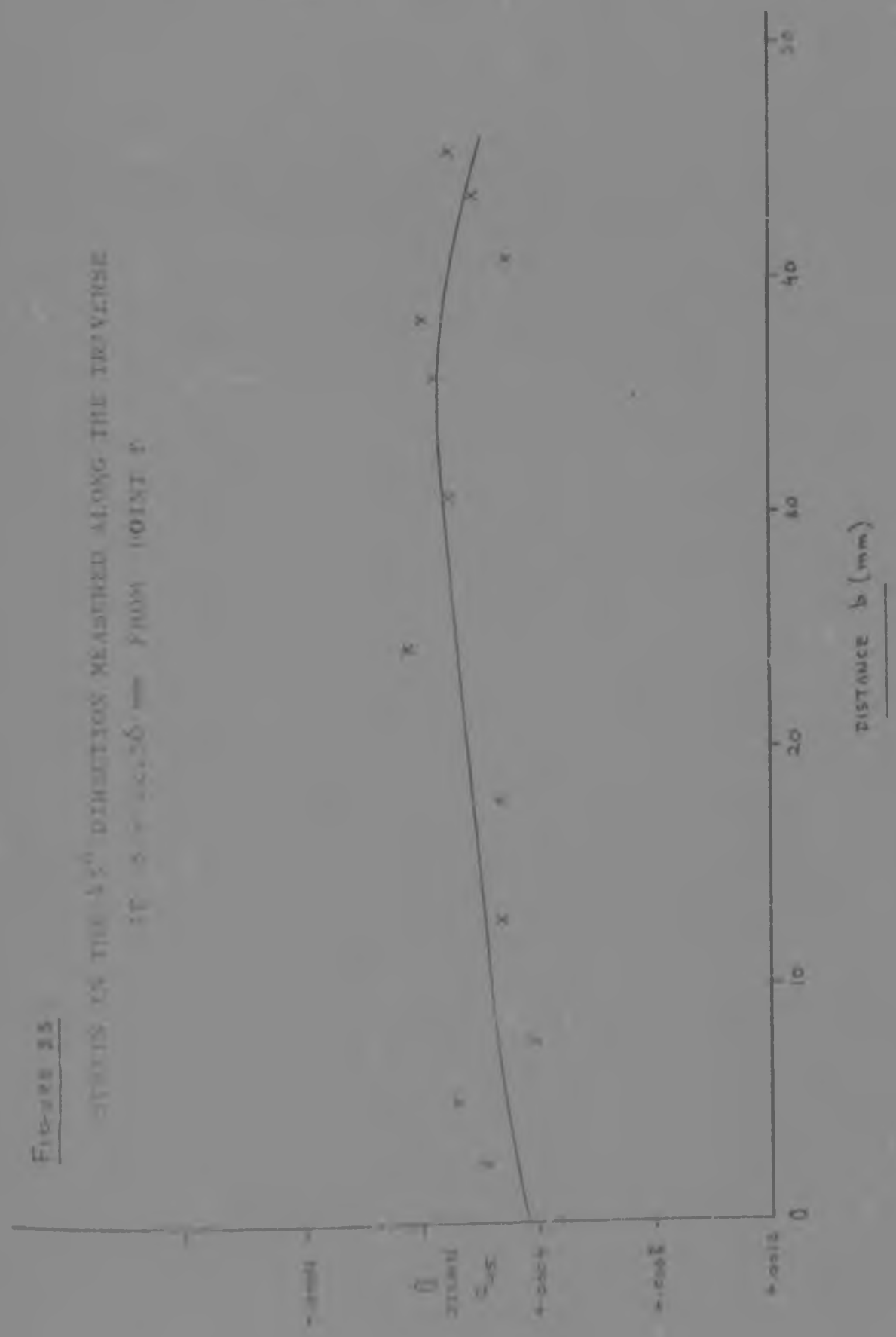


Figure 54.

Activity in the 15th population measured along the traverse
 $\Delta T = 5$ to 20,000 mm from point P

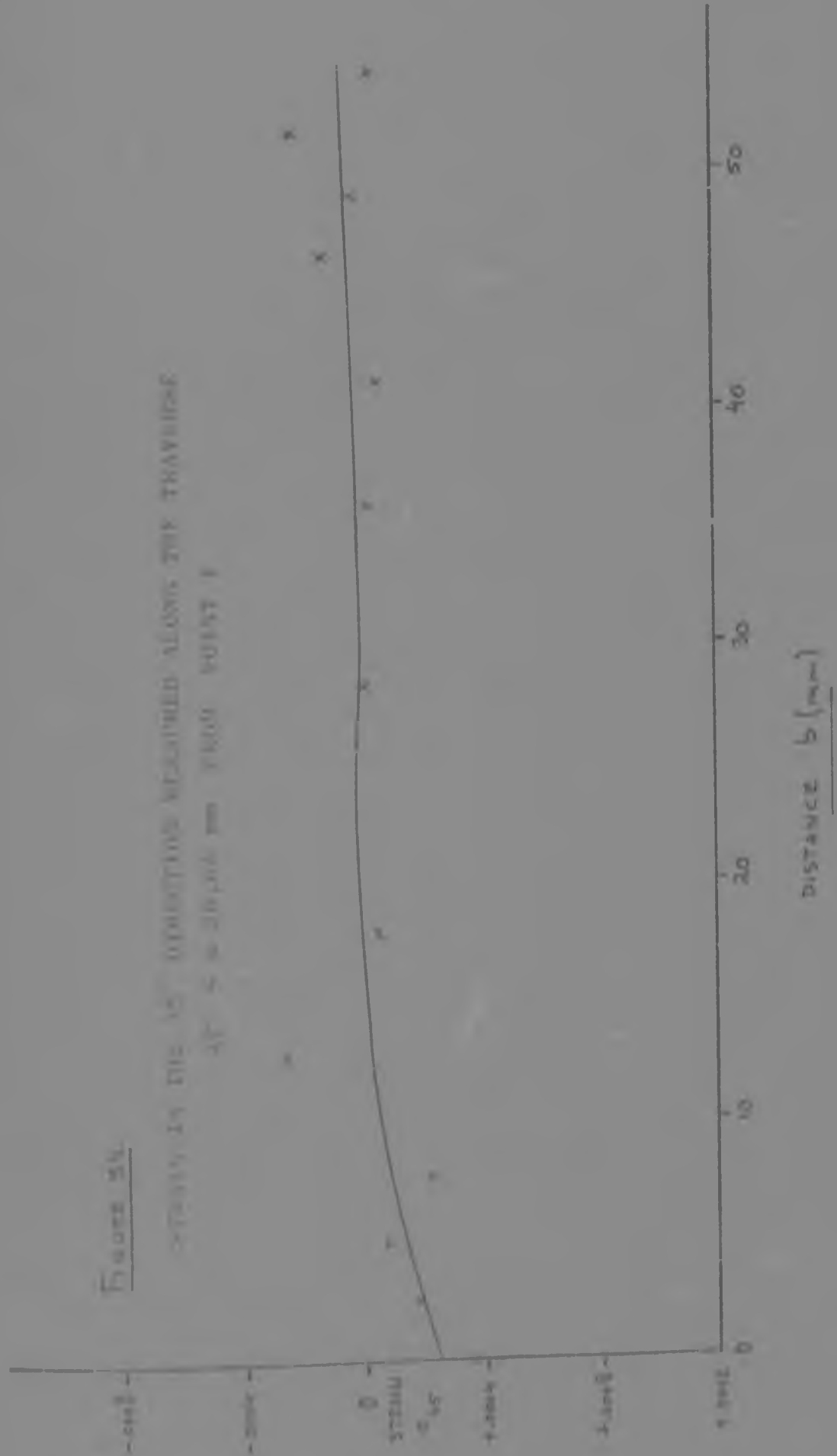


Figure 35.

Scatter in the y-direction searched along the traverse.

$$\Delta x = 2.5 \text{ mm}$$

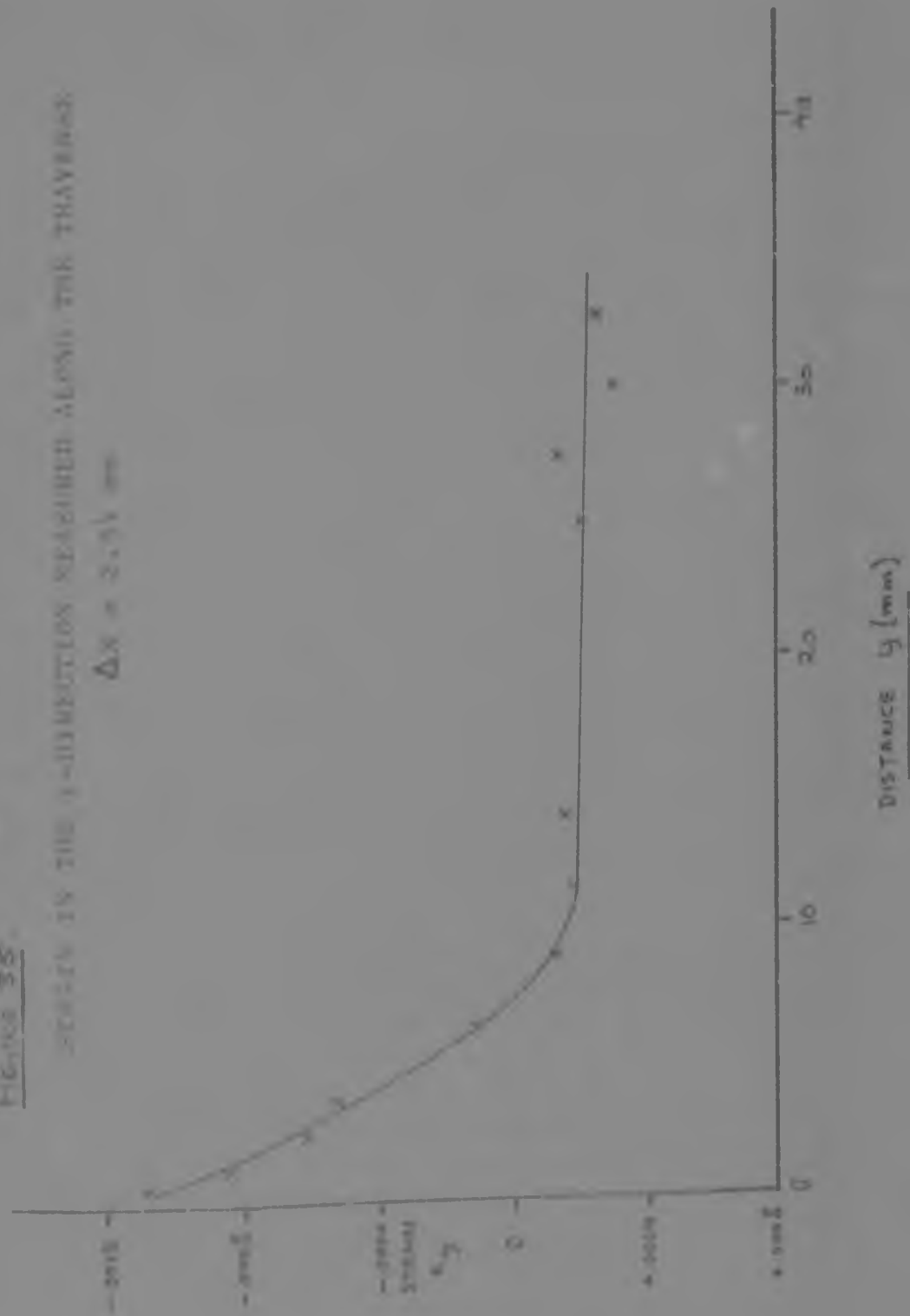


Figure 34.

DATA IS THE SUBSTITUTION OF AROUND ALONG THE TRAVERSE

$$\Delta x = 5.81 \text{ mm}$$

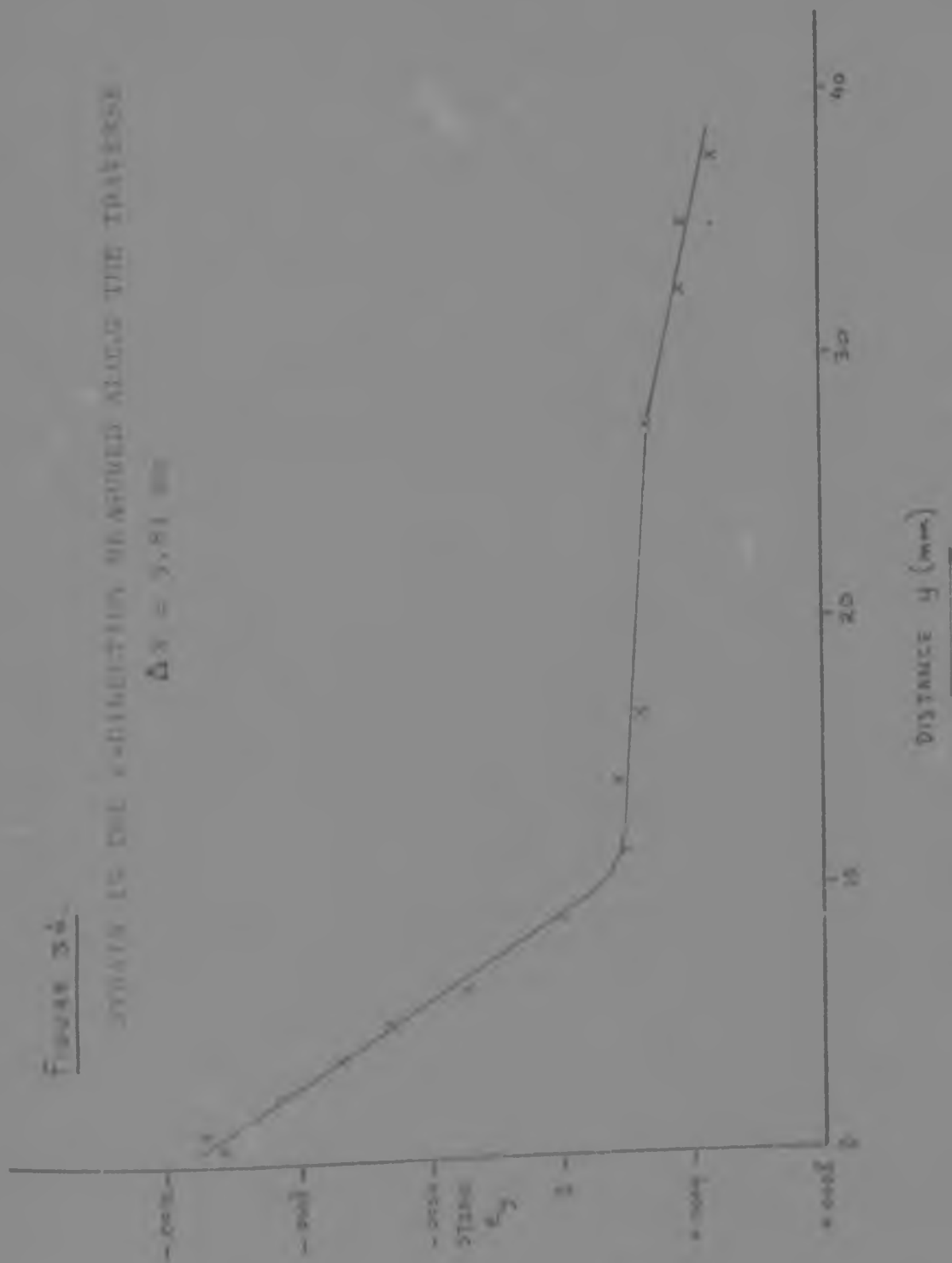


Figure 37

STRAIN IN THE x -DIRECTION MEASURED ALONG THE TRAVERSE

$$\Delta x = 0.10 \text{ mm}$$

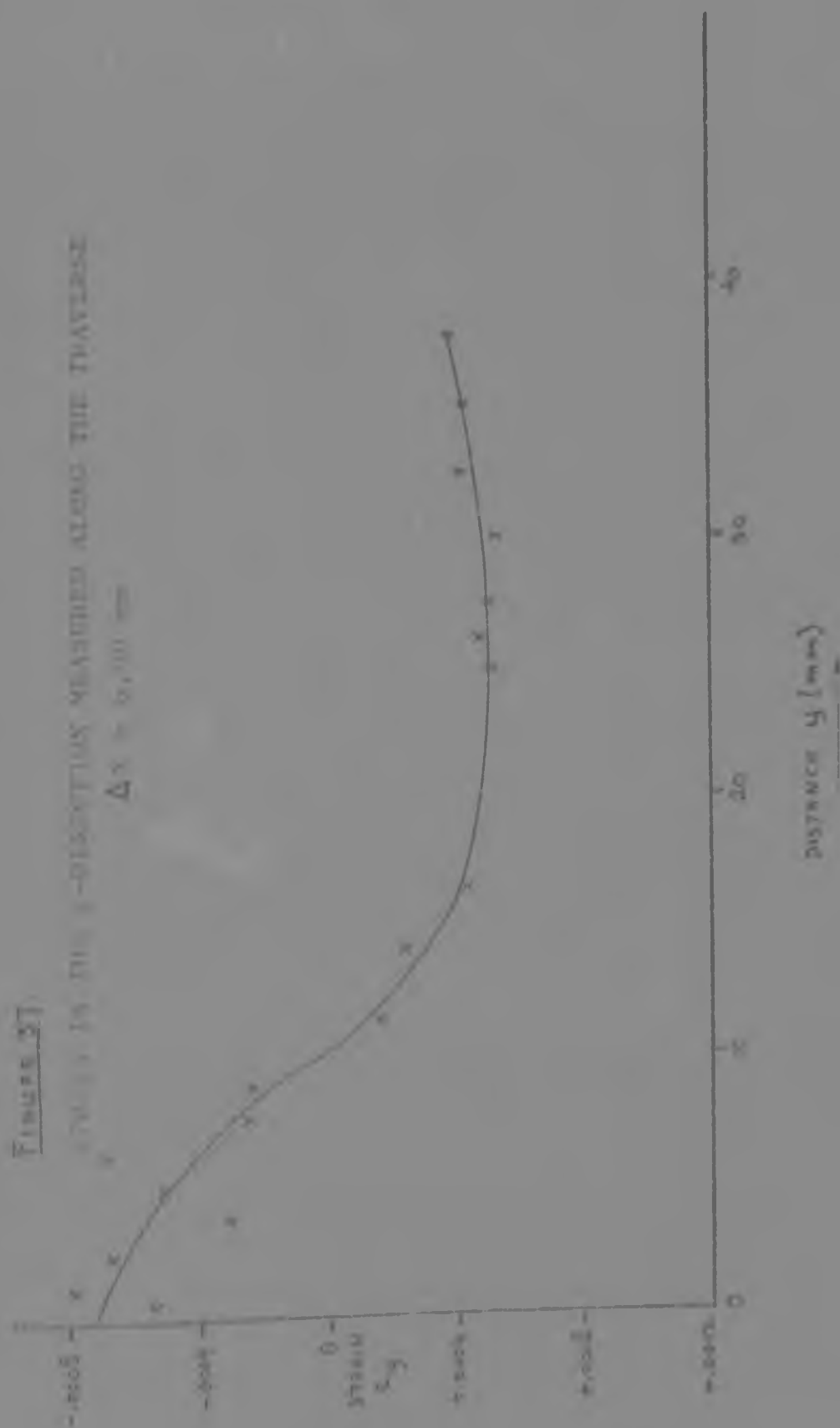


Figure 38: $\Delta x = 10.16 \text{ mm}$
 Figure 39: $\Delta x = 13.32 \text{ mm}$

Figure 38

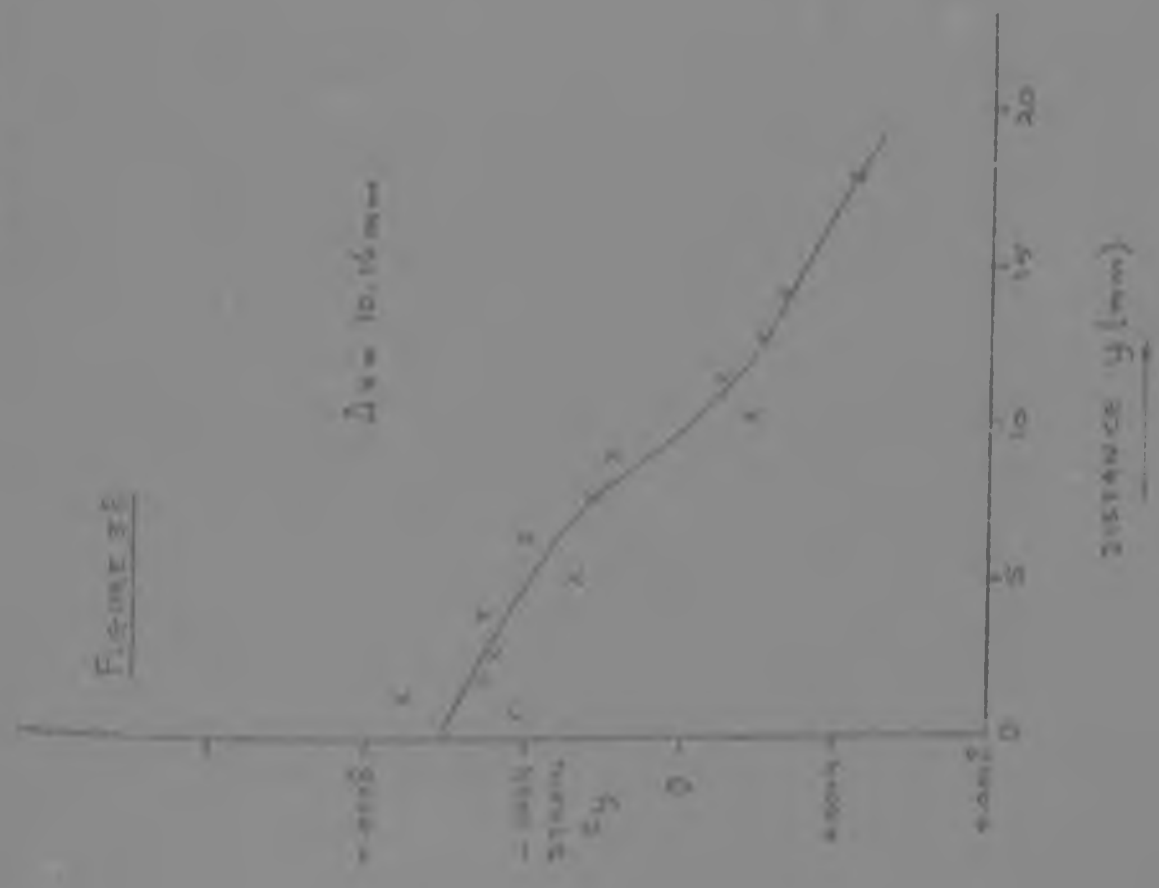


Figure 39

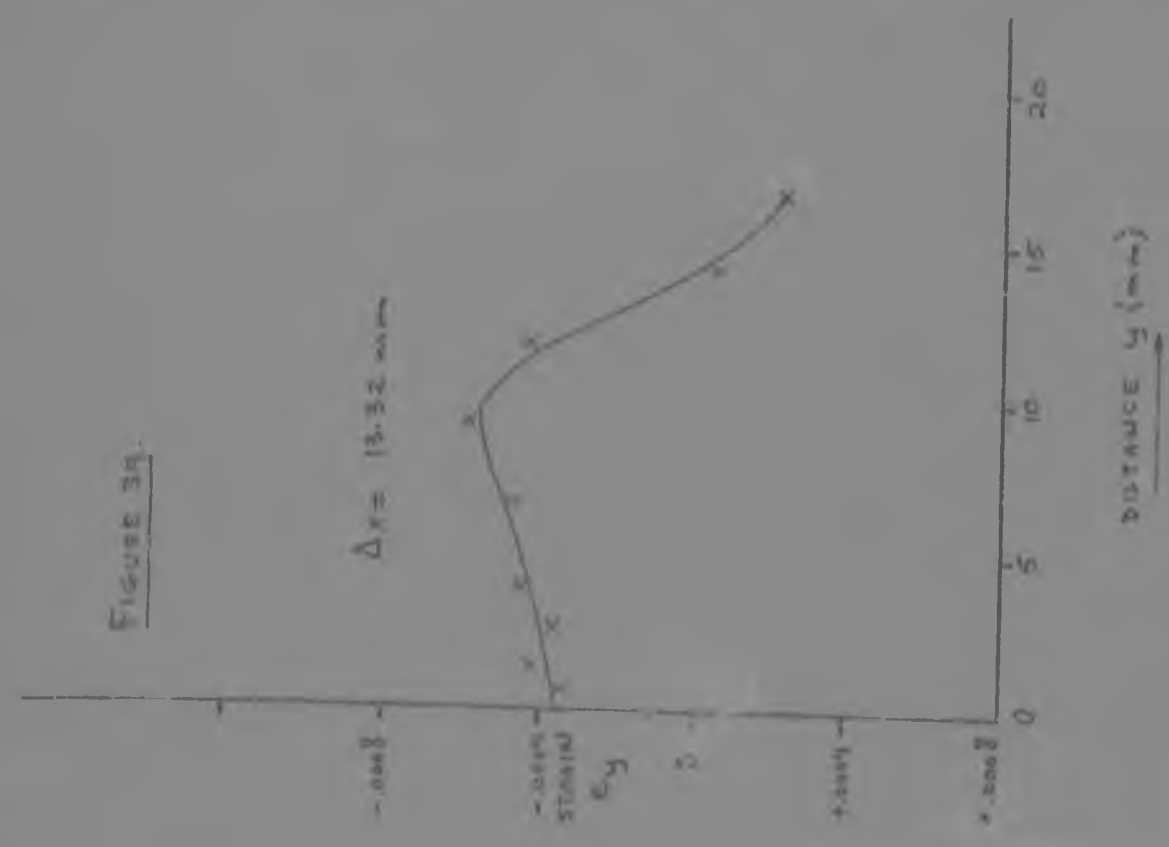
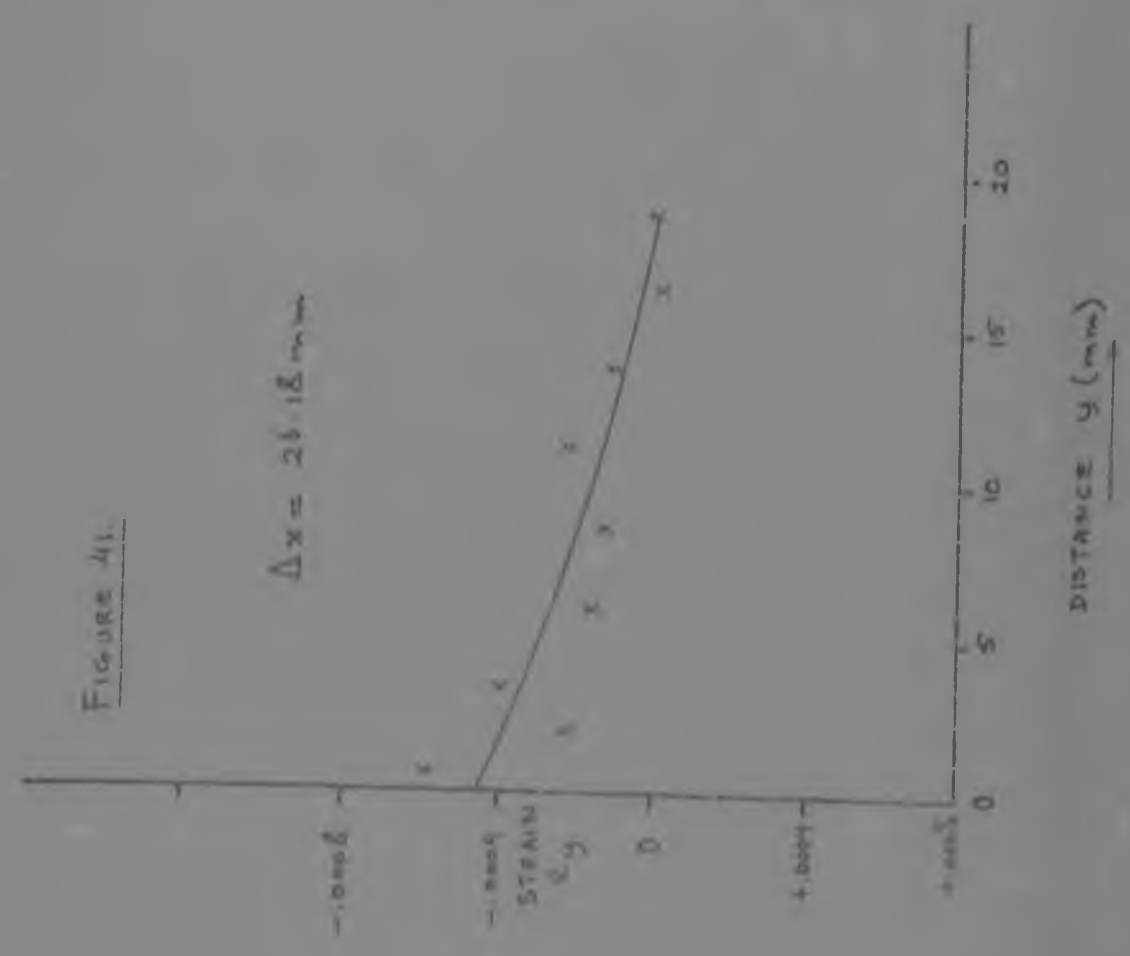
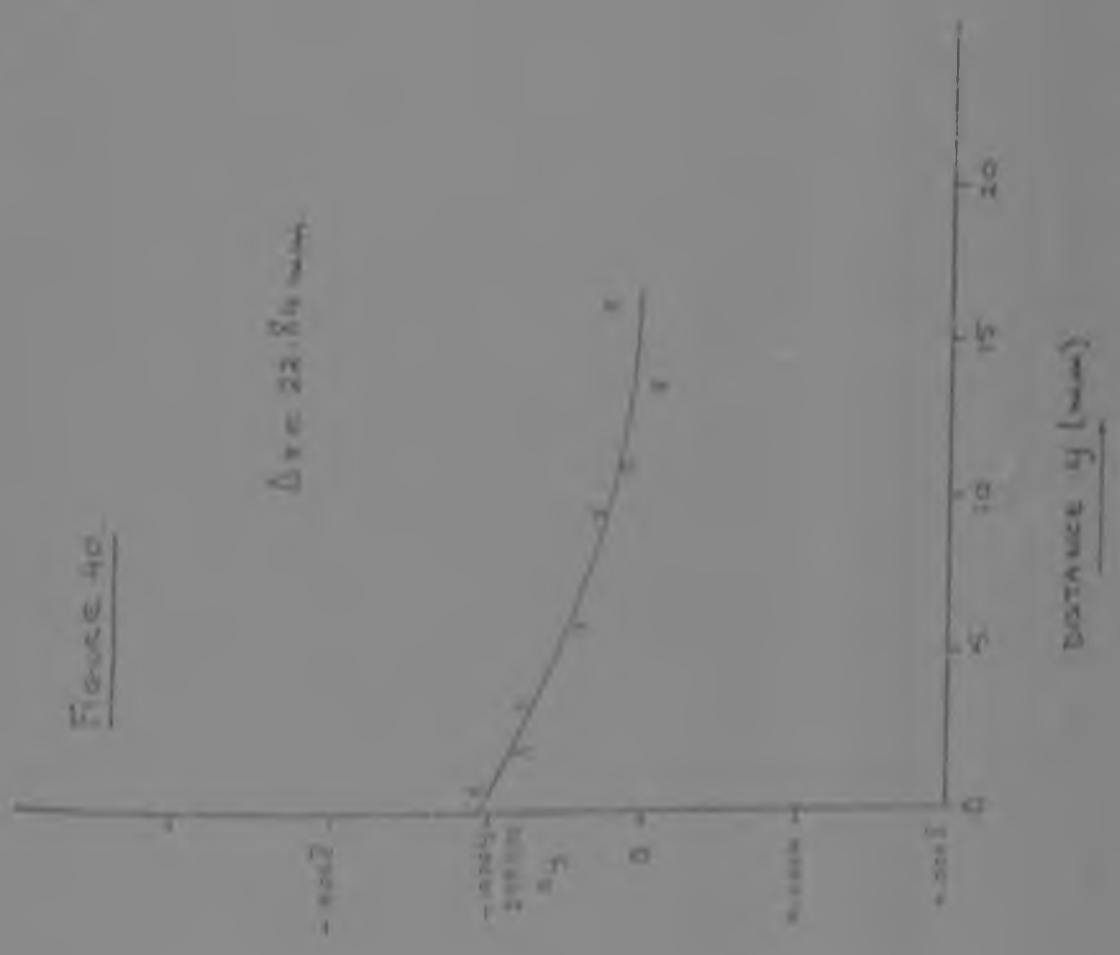


Figure 40 and Figure 41 show the relationship between the measured strain Δx and the distance y from the neutral axis.



DATA IN THE x-DIRECTION IS SHOWN ABOVE AND PRACTICALLY

$$\Delta x = 31.25 \text{ mm} \quad \text{AND} \quad \Delta x = 31.25 \text{ mm}$$

FIGURE 44

$$\Delta x = 31.25 \text{ mm}$$

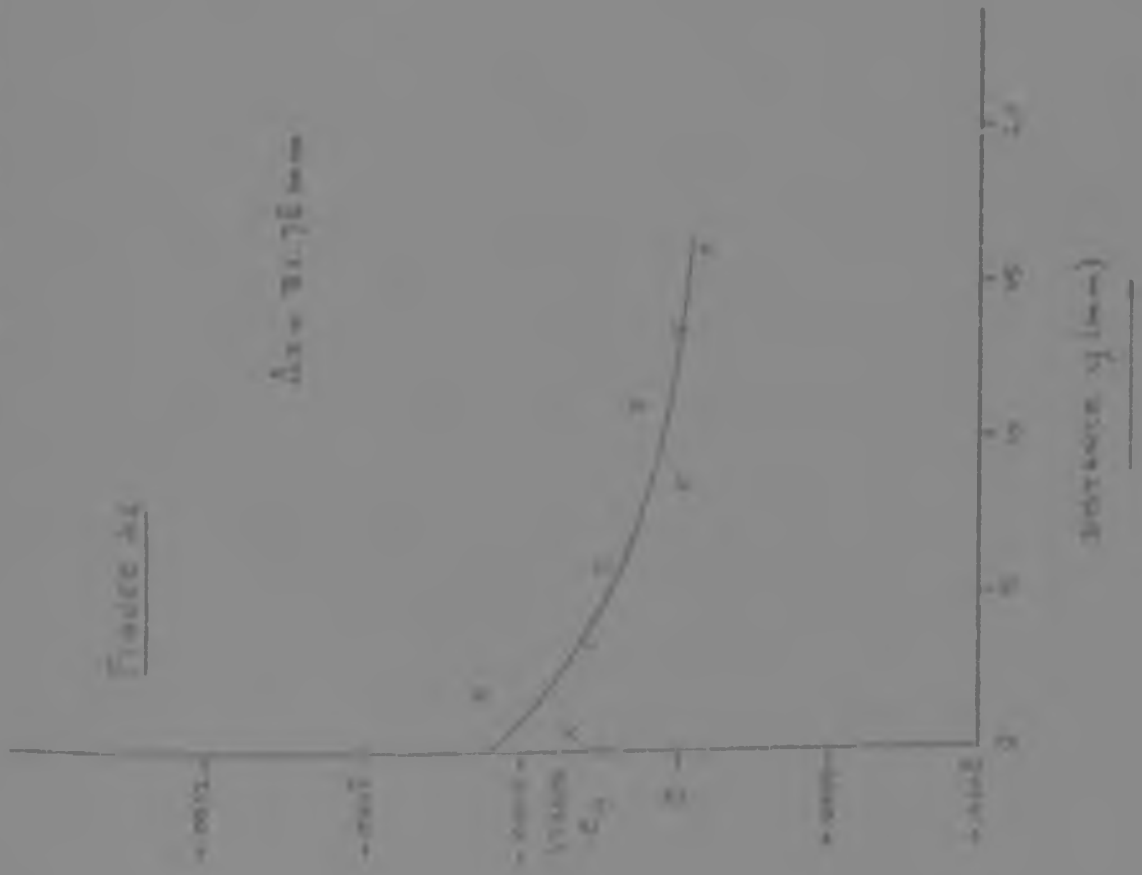


FIGURE 45

$$\Delta x = 31.25 \text{ mm}$$

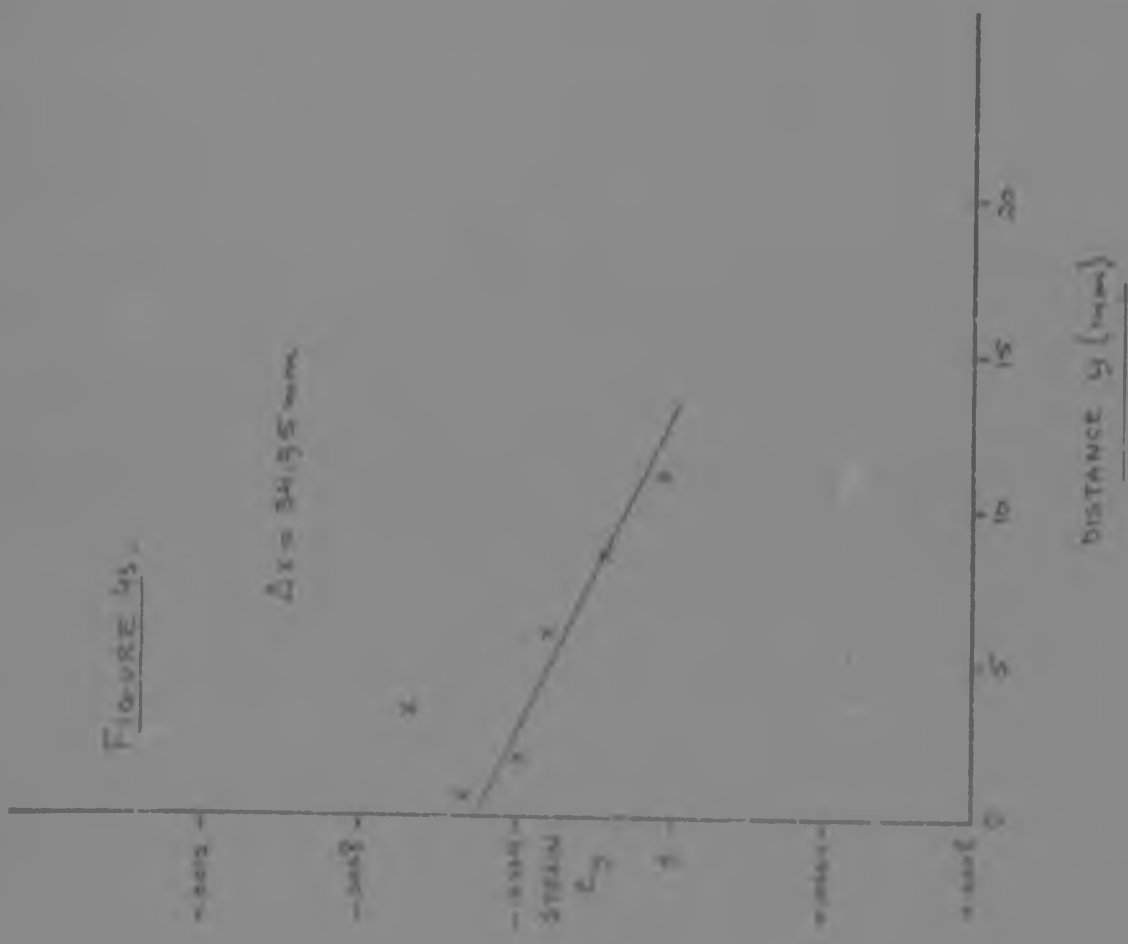


Figure 4a

STATUS IN THE Y-DIRECTION PLOTTED ALONG Δy TRAVERSES

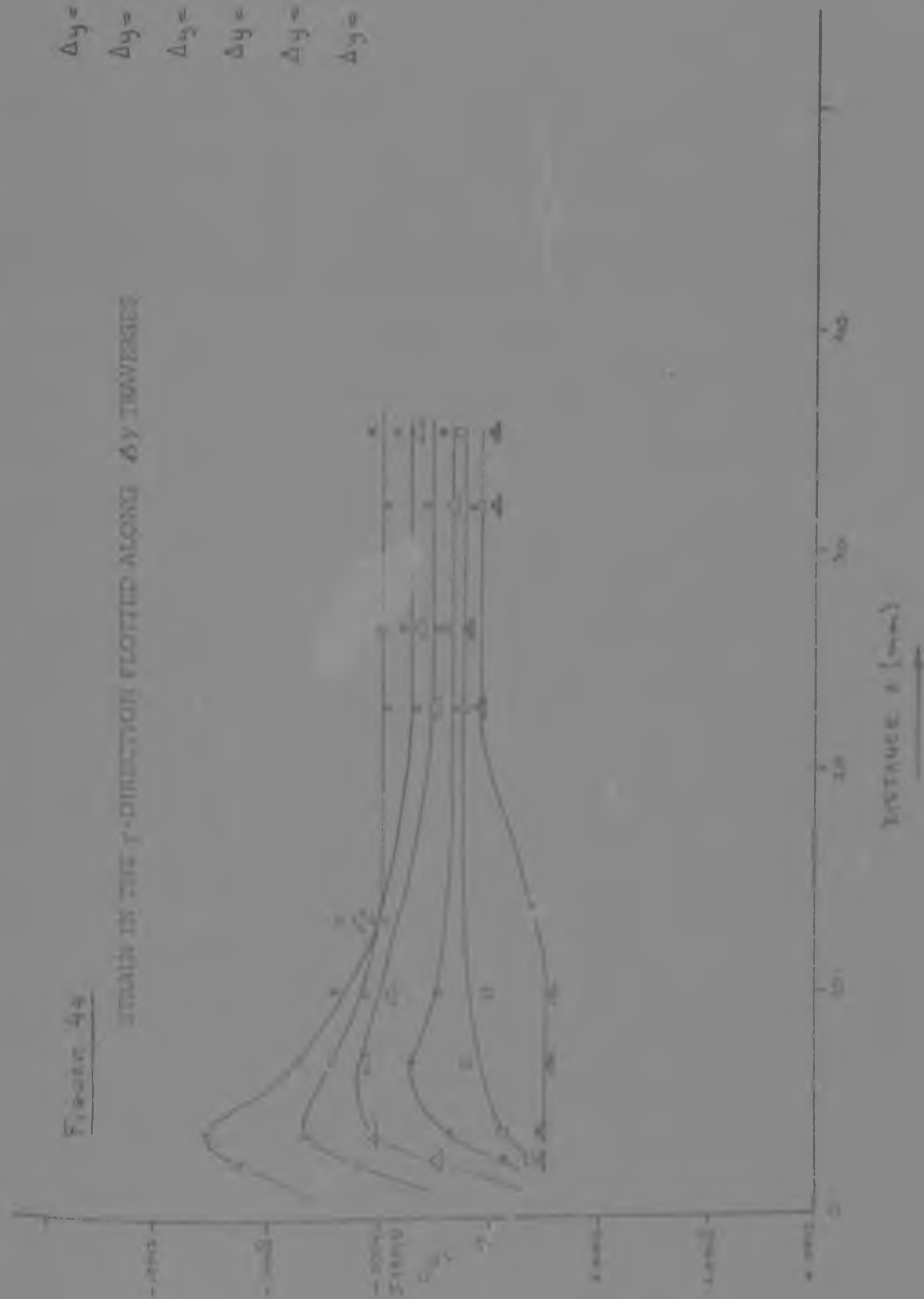
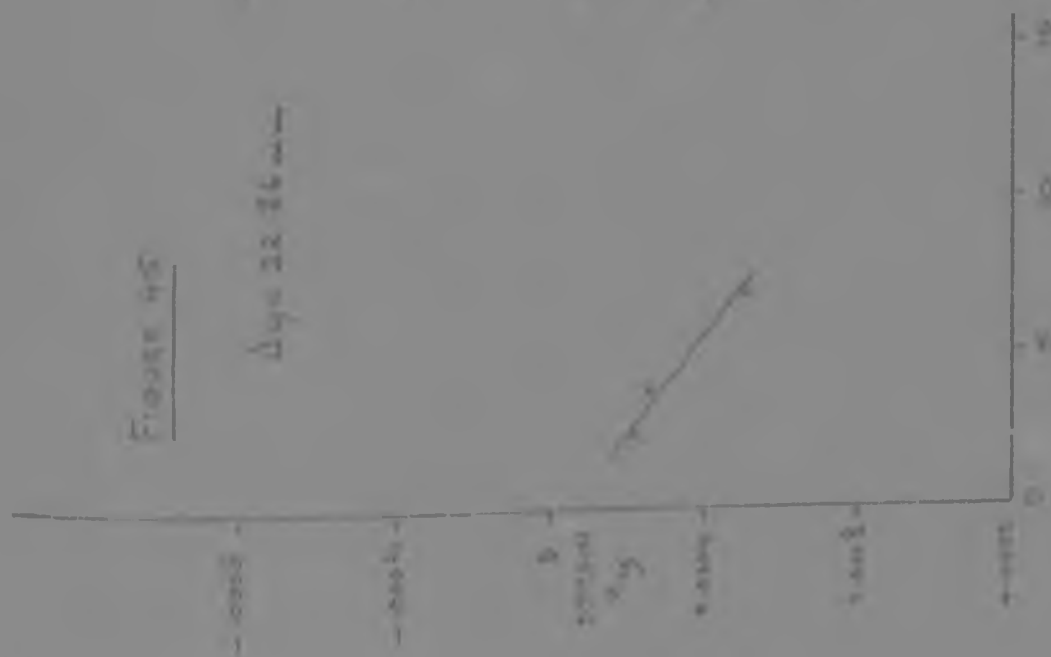


Figure 45, 46, 47. $\Delta y = 22.86 \text{ mm}$, $\Delta y = 27.94 \text{ mm}$, $\Delta y = 31.75 \text{ mm}$

Figure 45

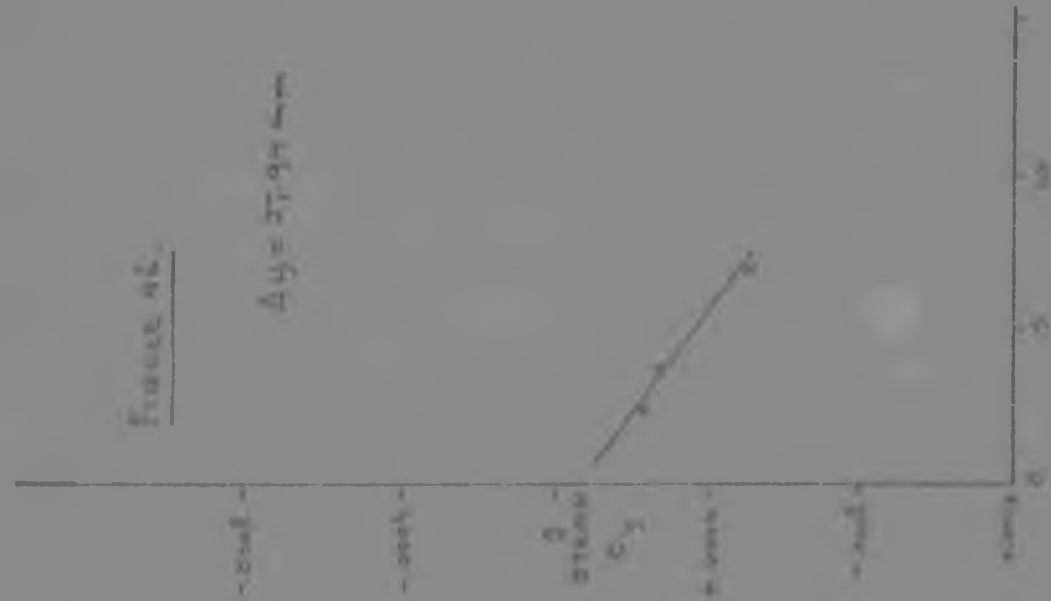
$\Delta y = 22.86 \text{ mm}$



DISTANCE x (mm)

Figure 46

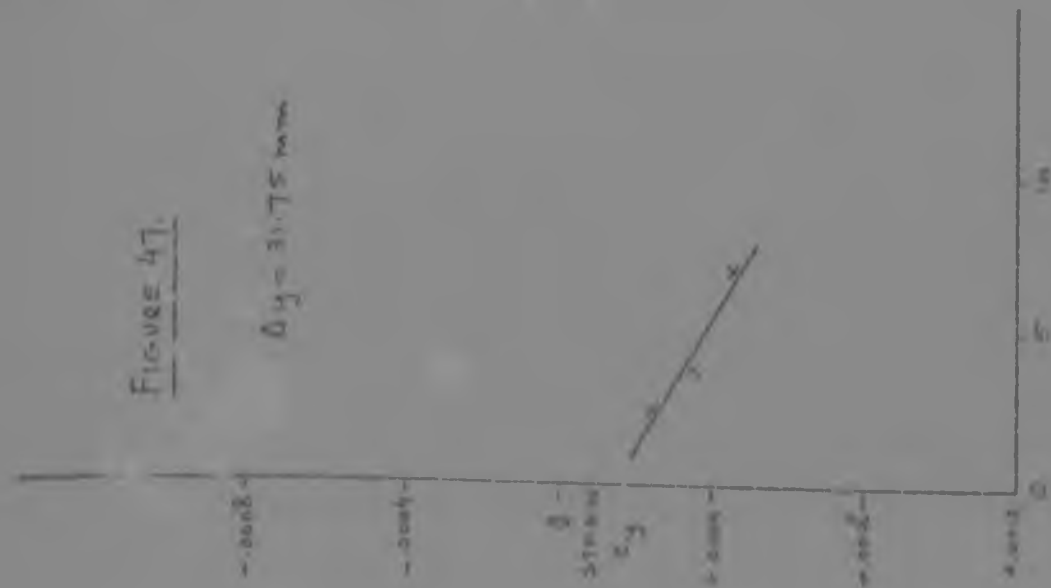
$\Delta y = 27.94 \text{ mm}$



DISTANCE x (mm)

Figure 47

$\Delta y = 31.75 \text{ mm}$



DISTANCE x (mm)

Figure A8. Paths in the x - y plane plotted along Δy traverses

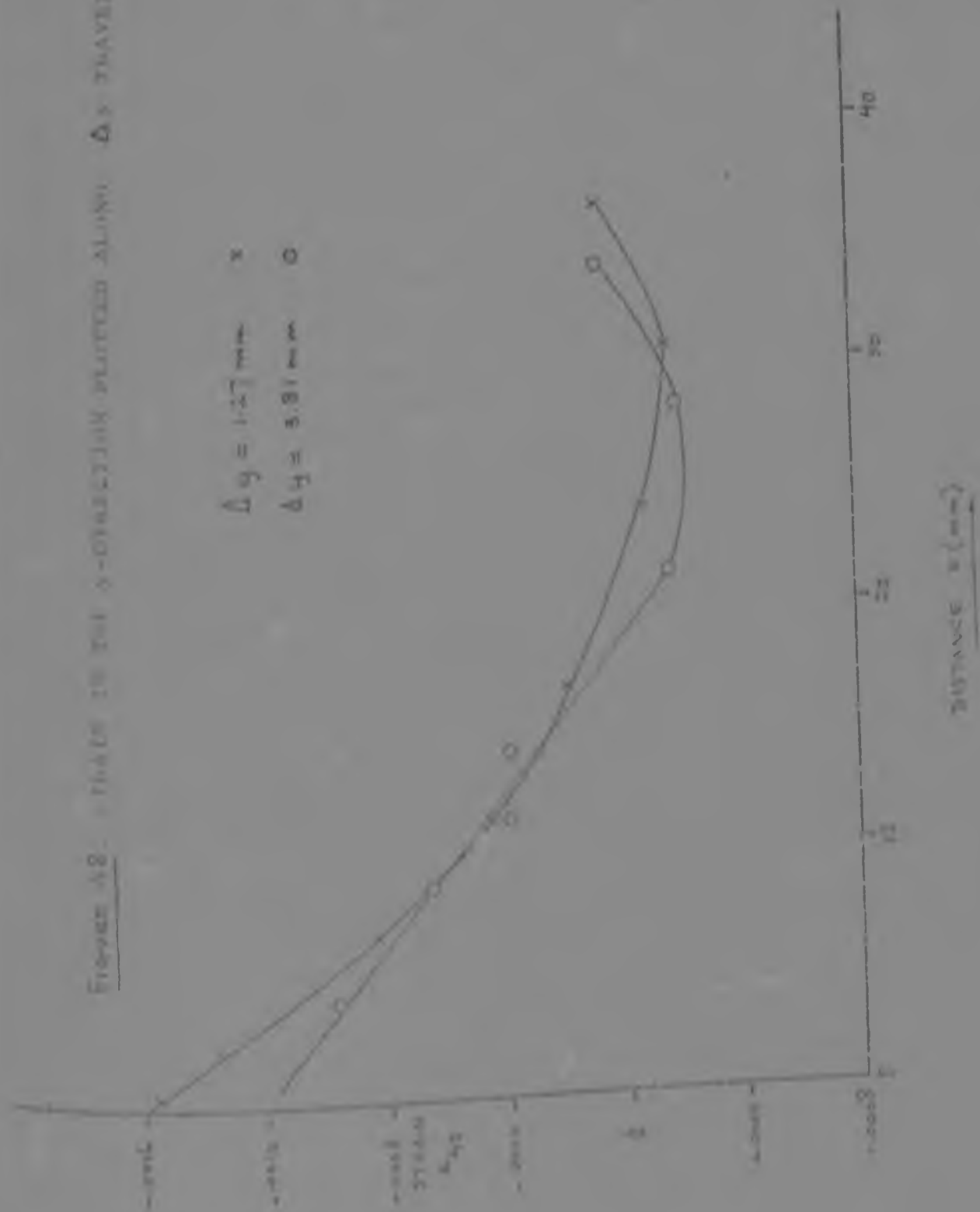


Figure 4a. ST-418 IN THE Δy DIAMETER PLOTTED ALONG Δy TRAVERSES

x $\Delta y = 5.71$ mm
 o $\Delta y = 8.00$ mm

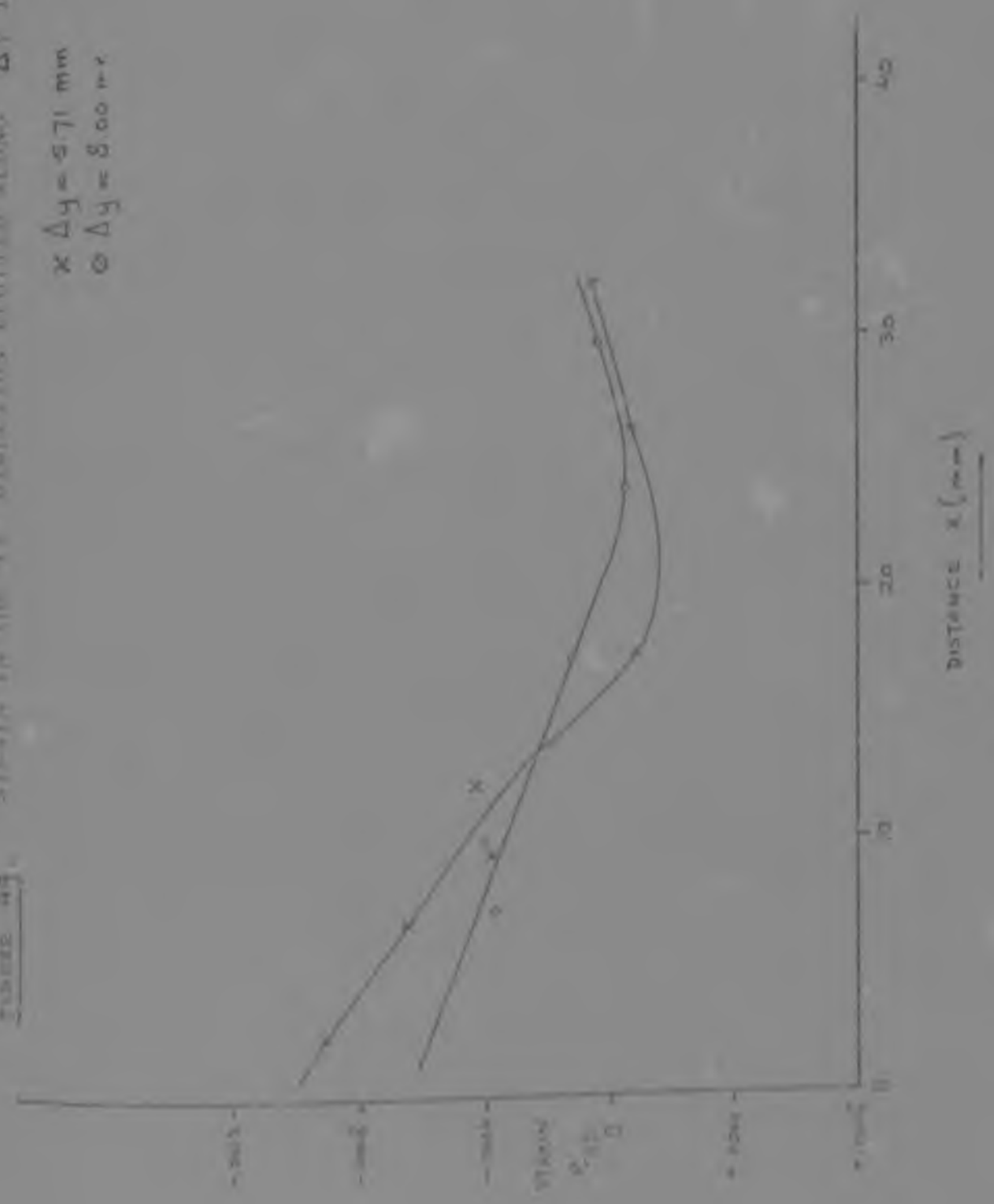


FIGURE 50 - STRAIN IN THE 90° DIRECTION LOCATED ALONG Δy TRAVERSES

$\Delta y = 9.58 \text{ mm}$ ϵ

$\Delta y = 13.10 \text{ mm}$ ϵ

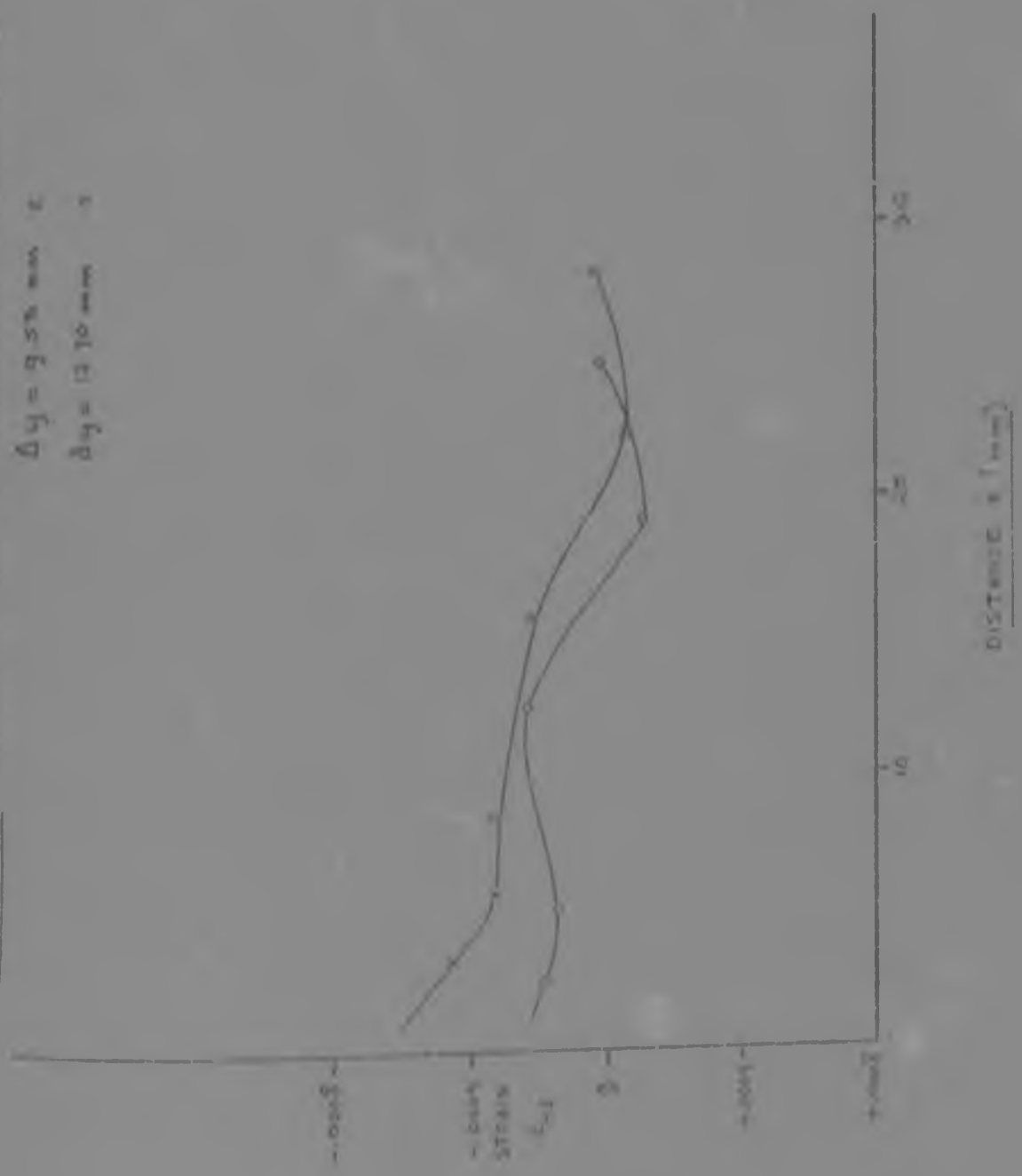


FIGURE 51 STRAIN IN THE 45° DIRECTION PLOTTED ALONG Δy TRAVERSES

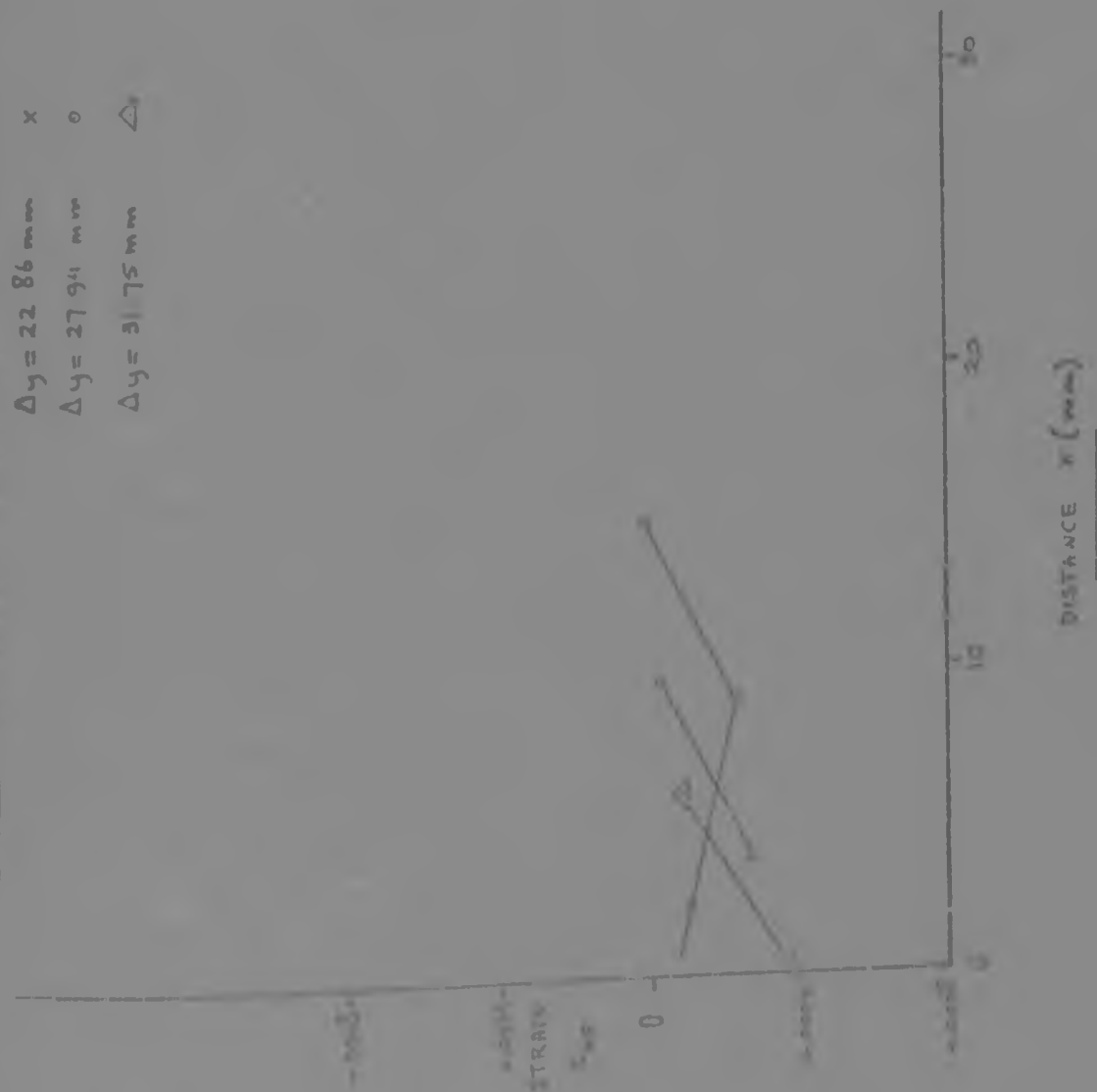


FIGURE 52. STAIN IN THE Y-DIRECTION PLOTTED ALONG ΔX TRAVERSES



Figure 5a. STRAIN IN THE X-DIRECTION PLOTTED ALONG ΔX TOLERANCES

$$\Delta y = 2.54 \text{ mm. } x$$

$$\Delta x = 6.99 \text{ mm. } 0$$

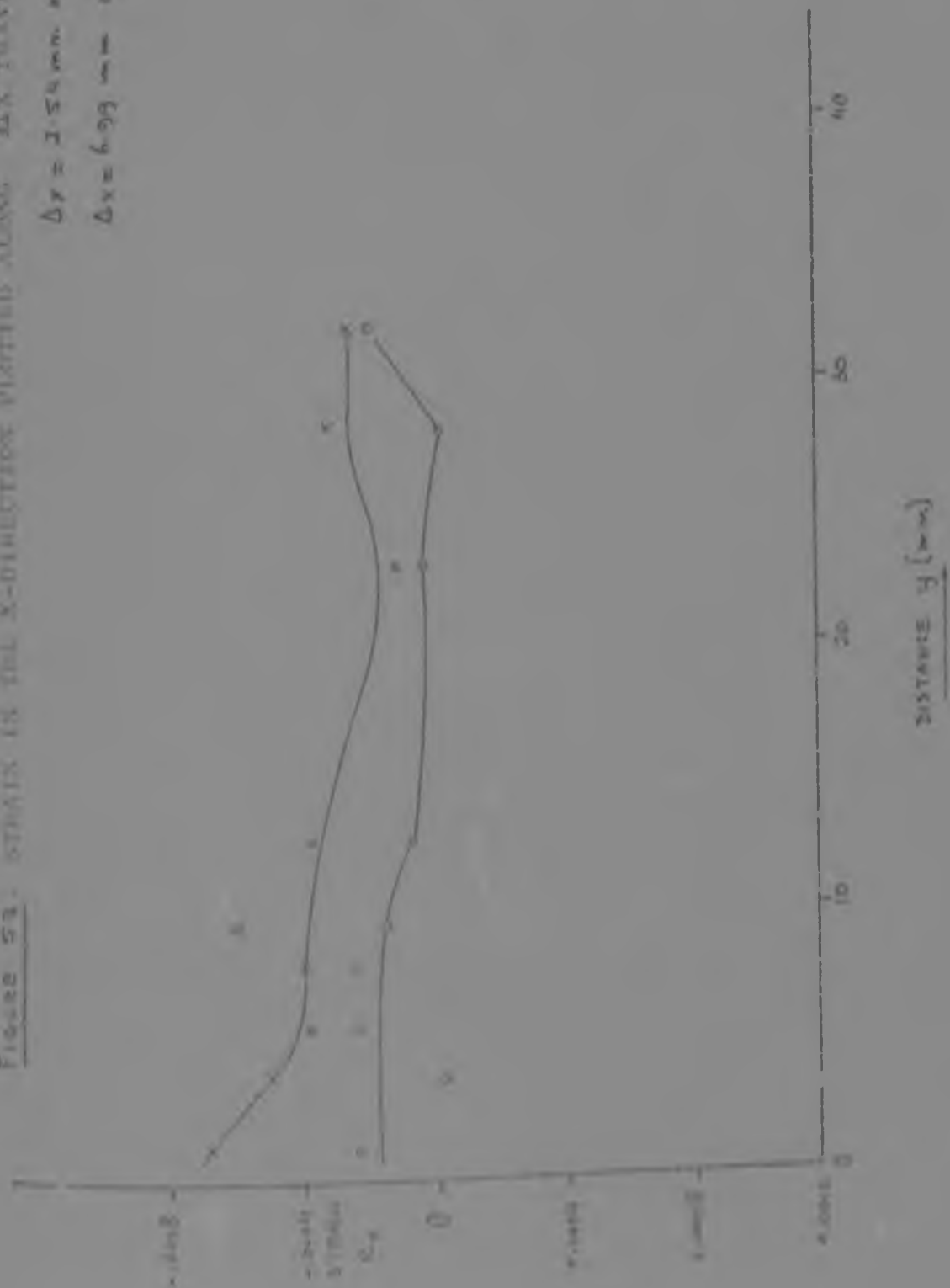


Figure 5a. Strain in the reduction plotted against distance along the traverse. $\Delta x = 10.16 \text{ mm}$ $\times$
 $\Delta x = 11.33 \text{ mm}$ $\circ$

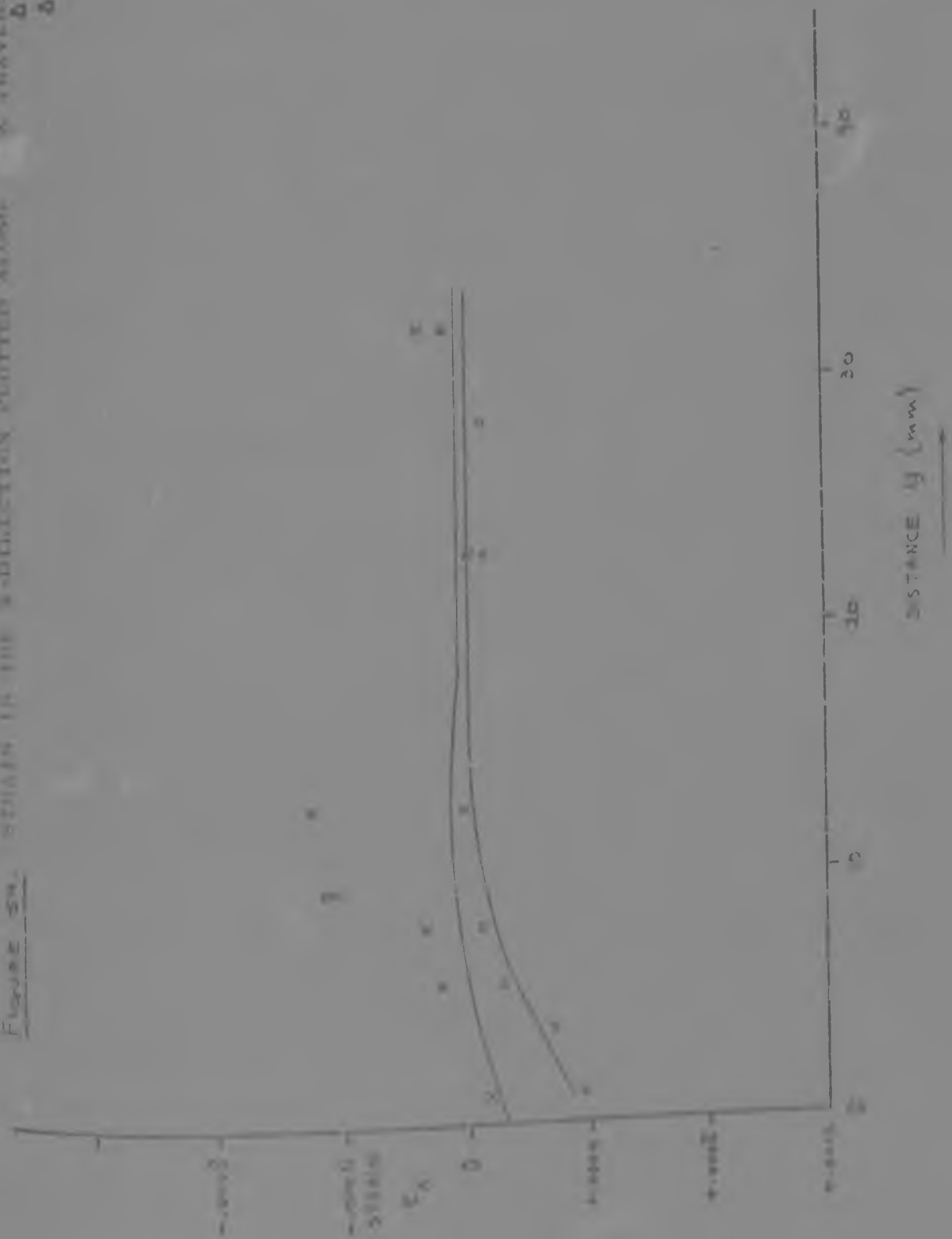


Figure 55.

Stress vs. distance from midspan for
 1000 lbs torque. $\Delta x = 22.16$ mm.
 $\Delta y = 26.16$ mm.

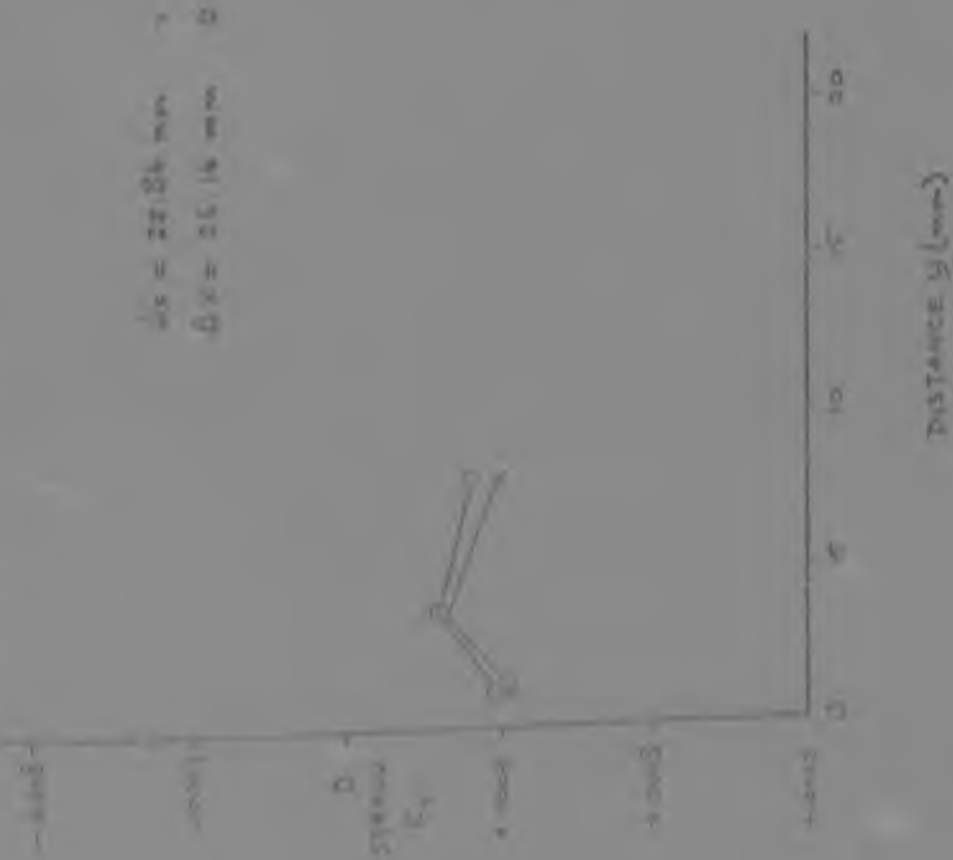


Figure 56.

Stress vs. distance from midspan for
 1000 lbs torque. $\Delta x = 31.78$ mm.

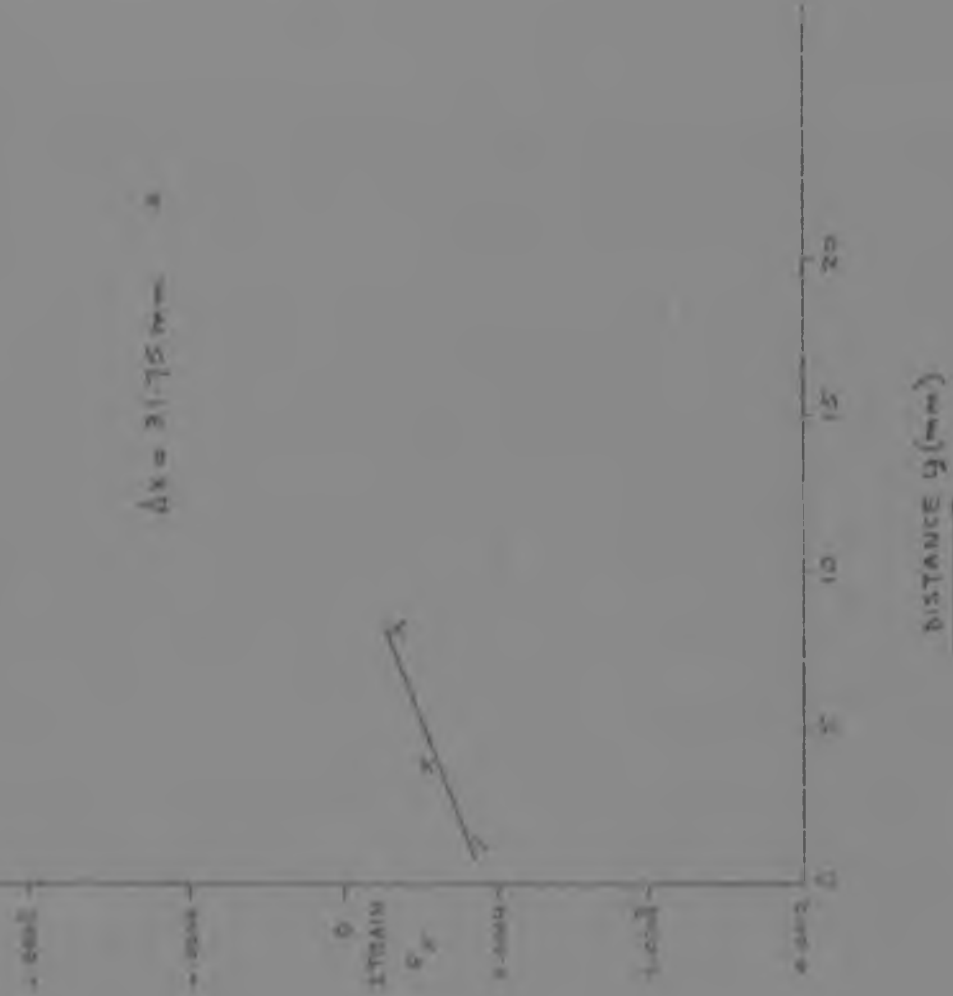


Figure 57. REFLECTION MEASURED ALONG Δx TRAVERSES

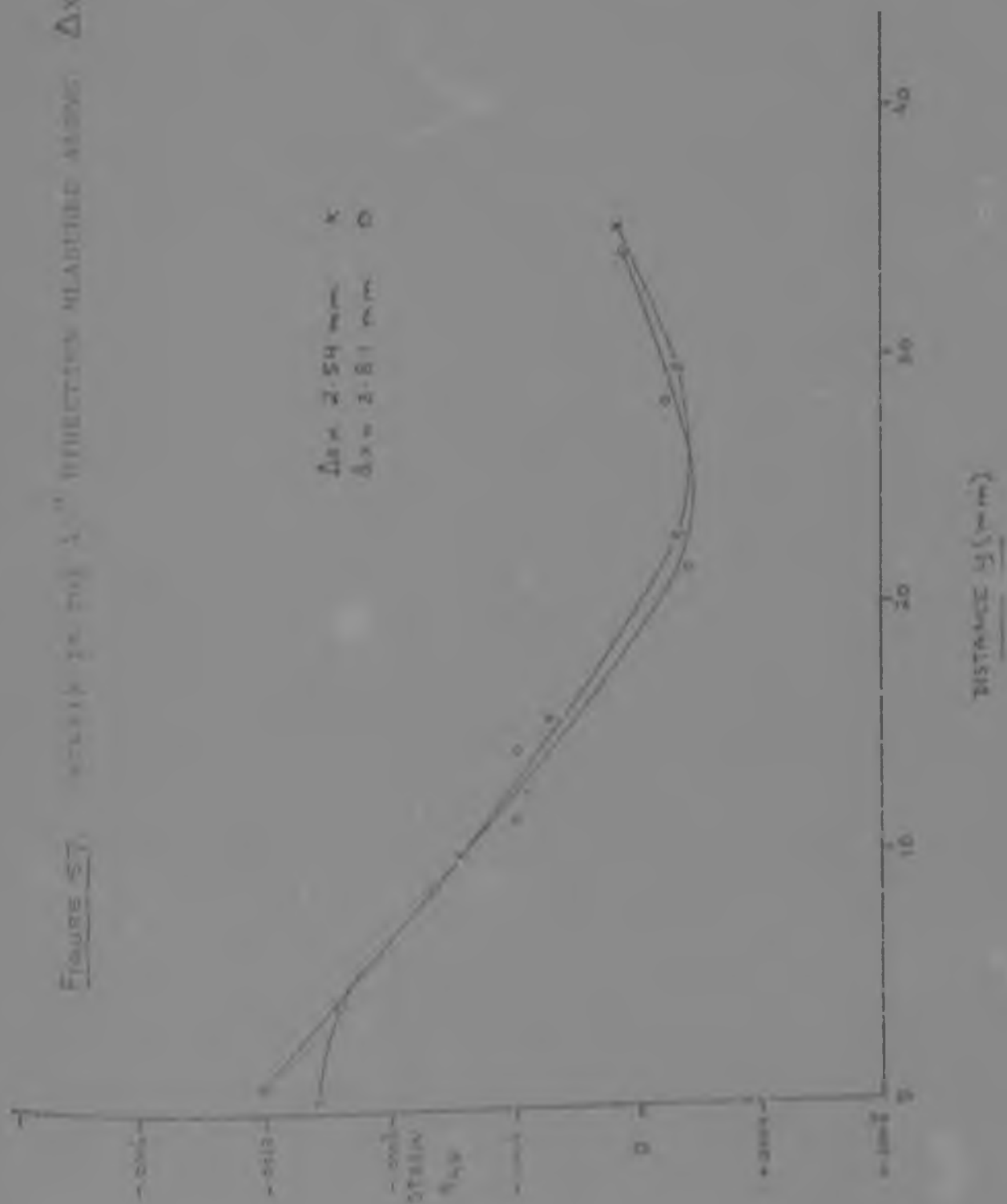
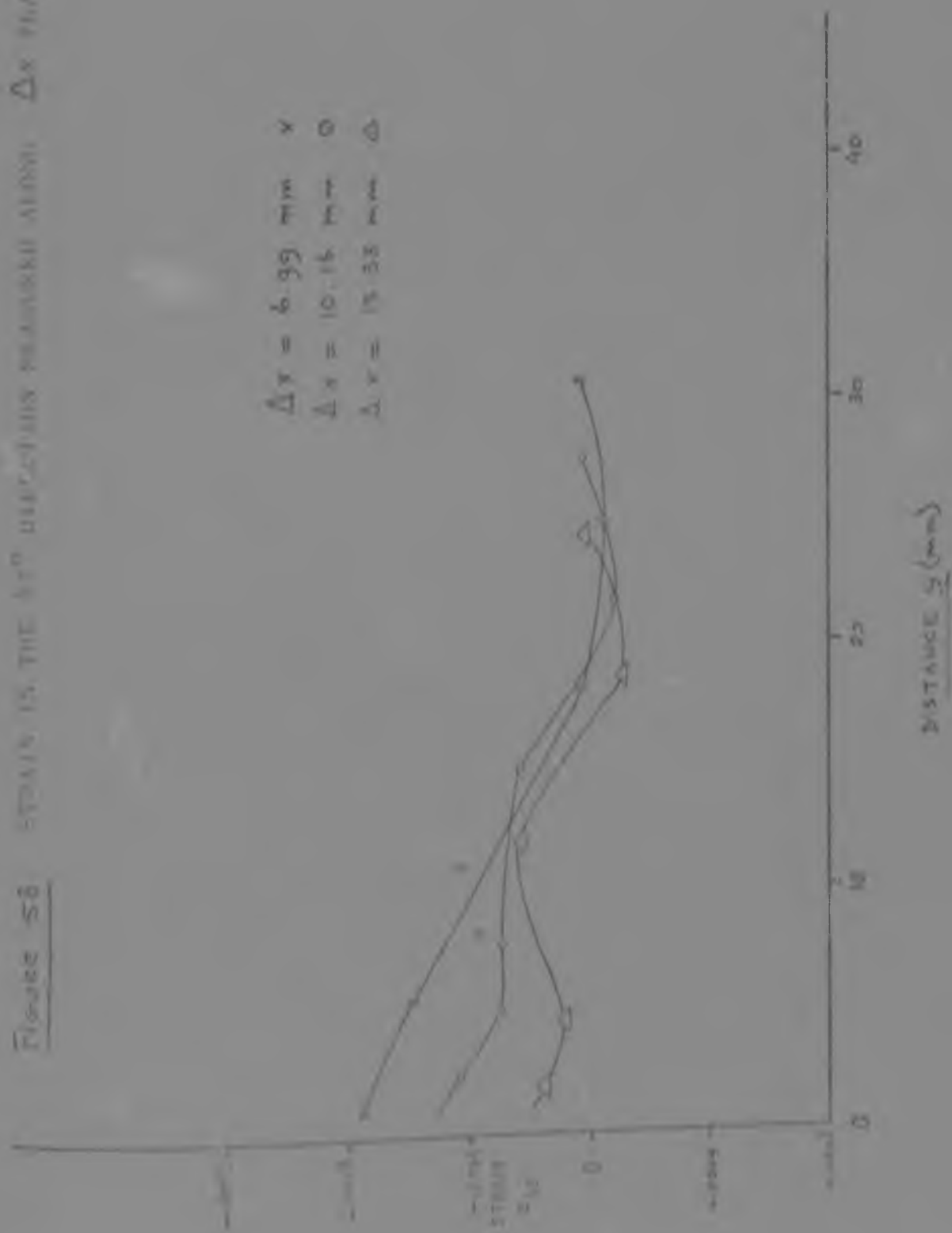
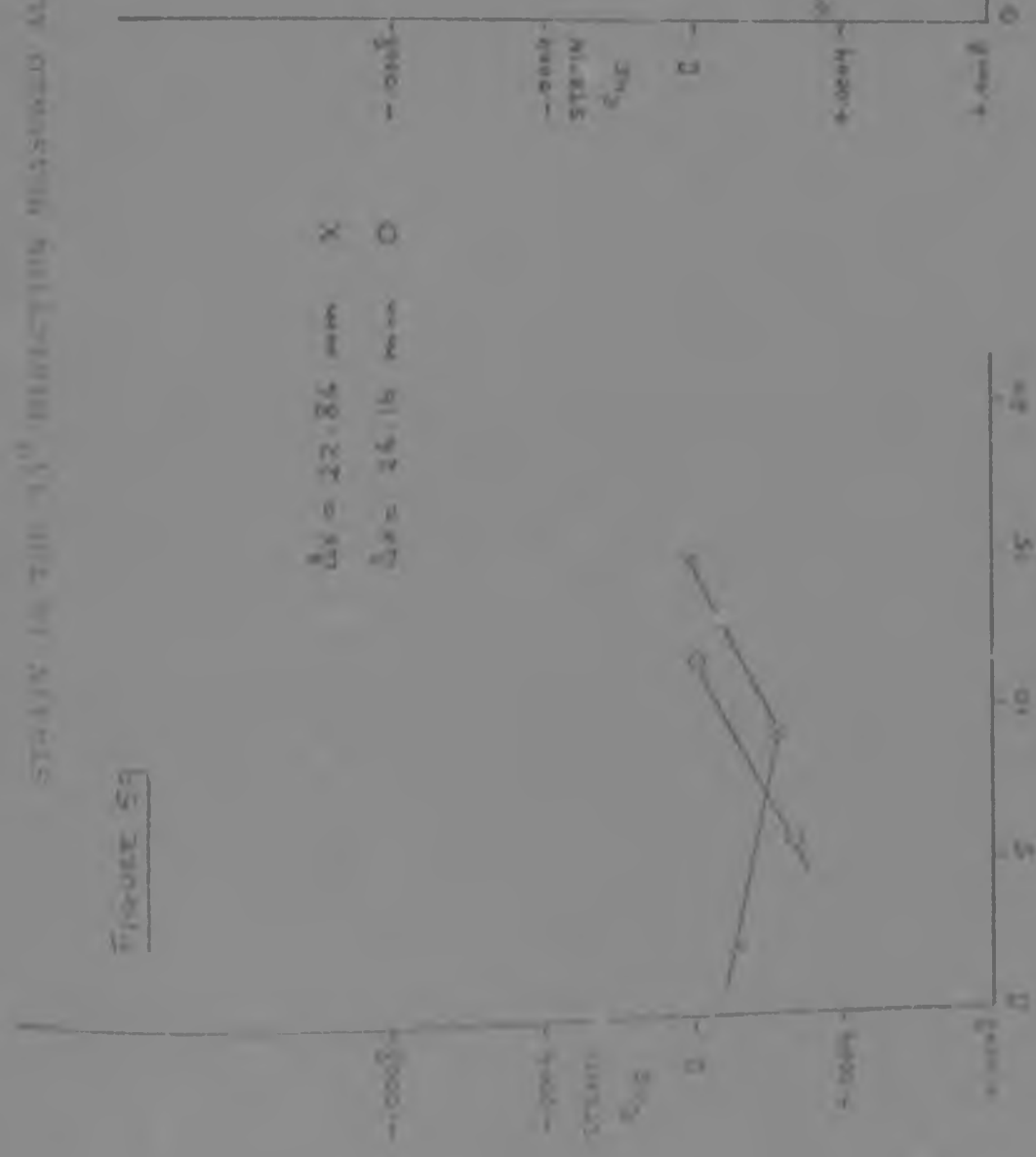


Figure 56 STRAIN IN THE 1st DEFLECTION MEASURED ALONG Δx FLANGES



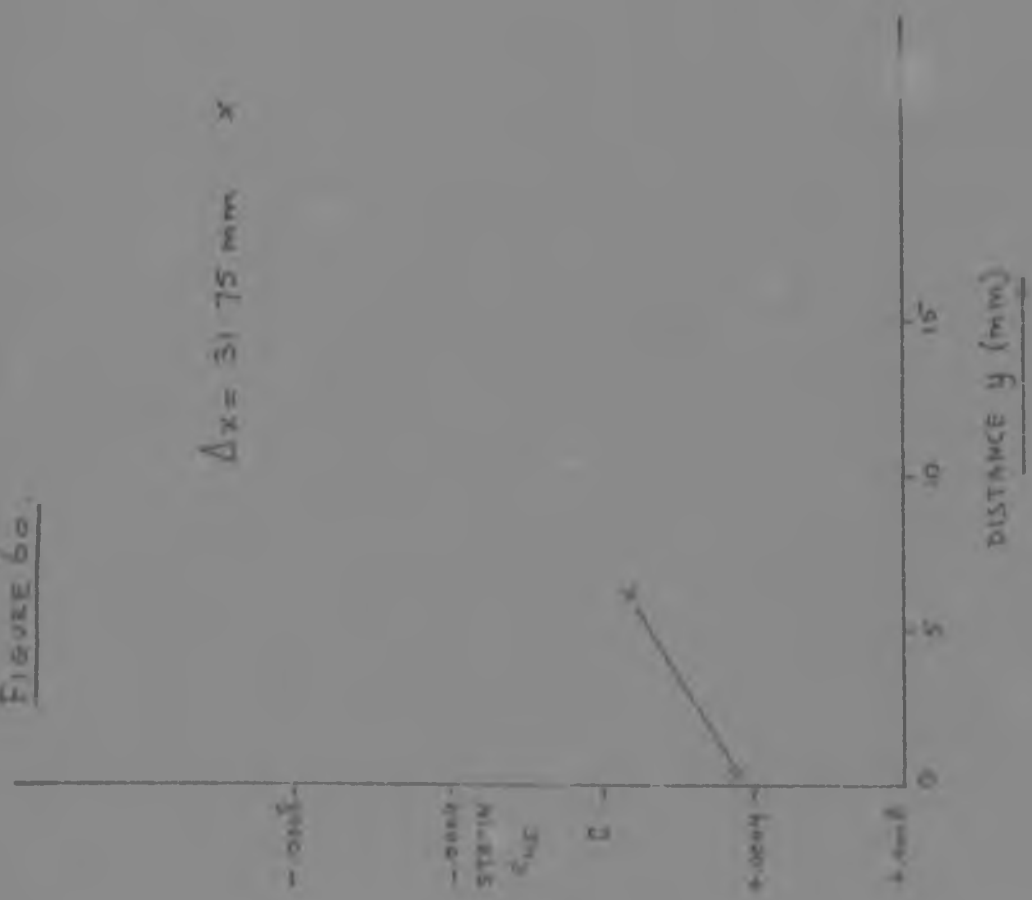
STRAIN IN THE 10^6 INJECTING MEASURED ALONG Δx TRAVERSES

FIGURE 59



$\Delta x = 22.86 \text{ mm}$ x
 $\Delta x = 26.16 \text{ mm}$ o

FIGURE 60



$\Delta x = 31.75 \text{ mm}$ x

Figure 61.

DISTRIBUTION OF THE σ_1 TENSILES ACROSS THE STEEL SURFACE
OF THE ANALOGUE (X-RAY MEASUREMENTS FROM TABLE XI)

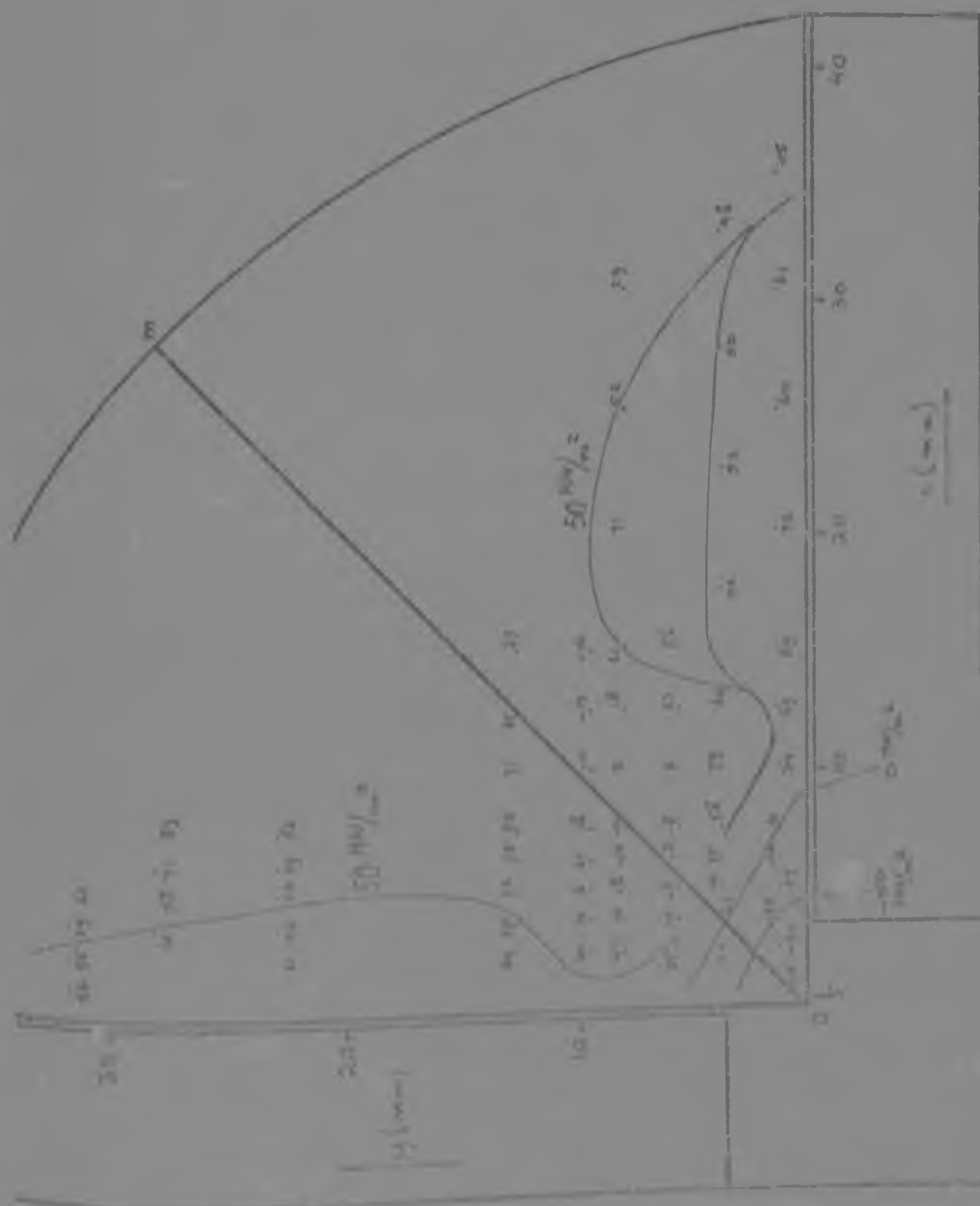


FIGURE 62

DISTRIBUTION OF THE σ_1 TENSILE STRESS ALONG THE STEEL SURFACE
OF THE ANALOGUE (X-RAY MEASUREMENTS FROM TABLE XI)



Figure 6a.

DISTRIBUTION OF THE Q_2 TUBES ACROSS THE SURFACE
OF THE ANALOGUE (X-RAY MEASUREMENTS FROM TABLE XI).

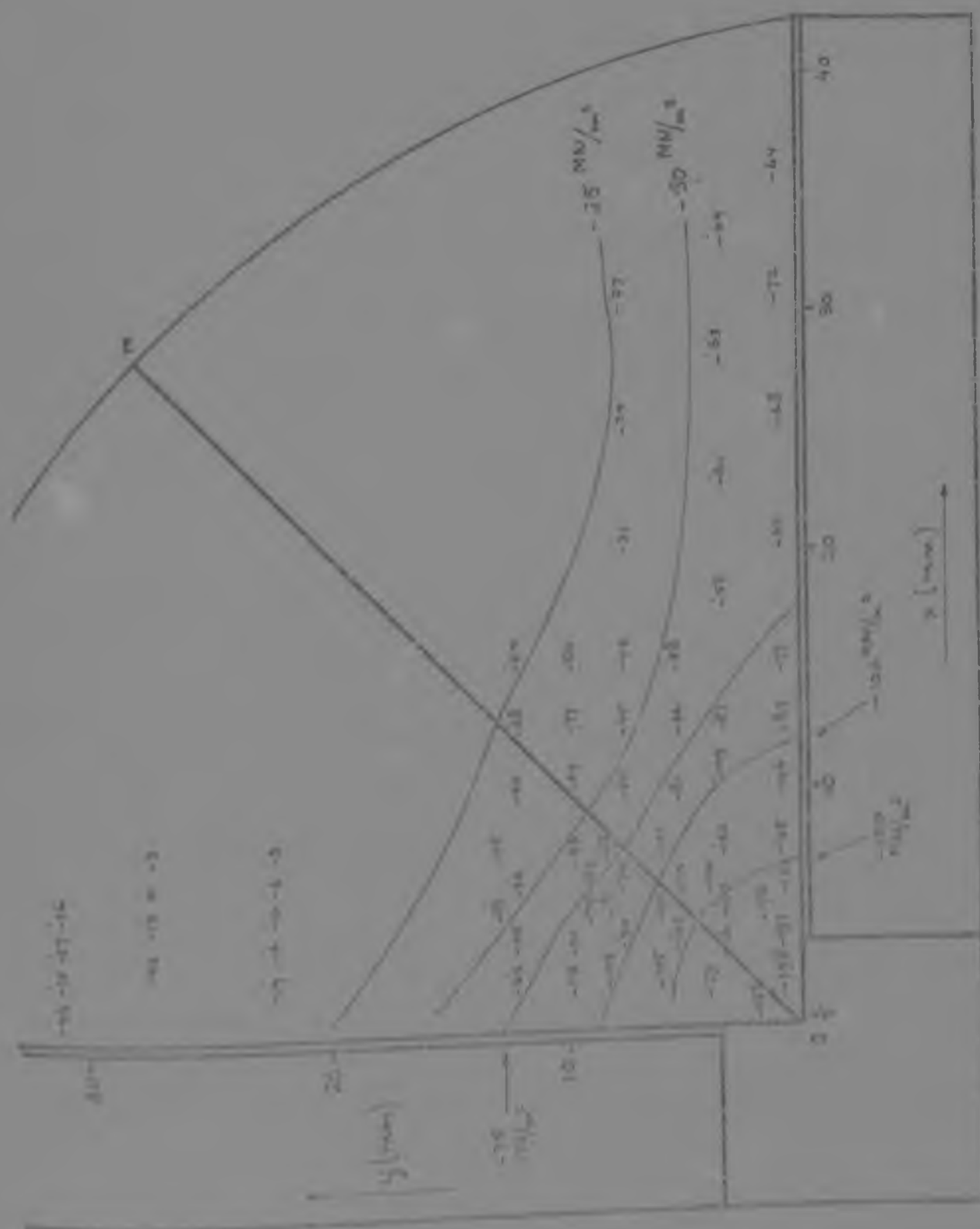


Figure 69.

DISTRIBUTION OF THE α_2 TEMPERATURE ALONG THE STEEL SURFACE
OF THE ANALOGUE (S-45Y 0.25% C STEEL FOR TABLE 21).

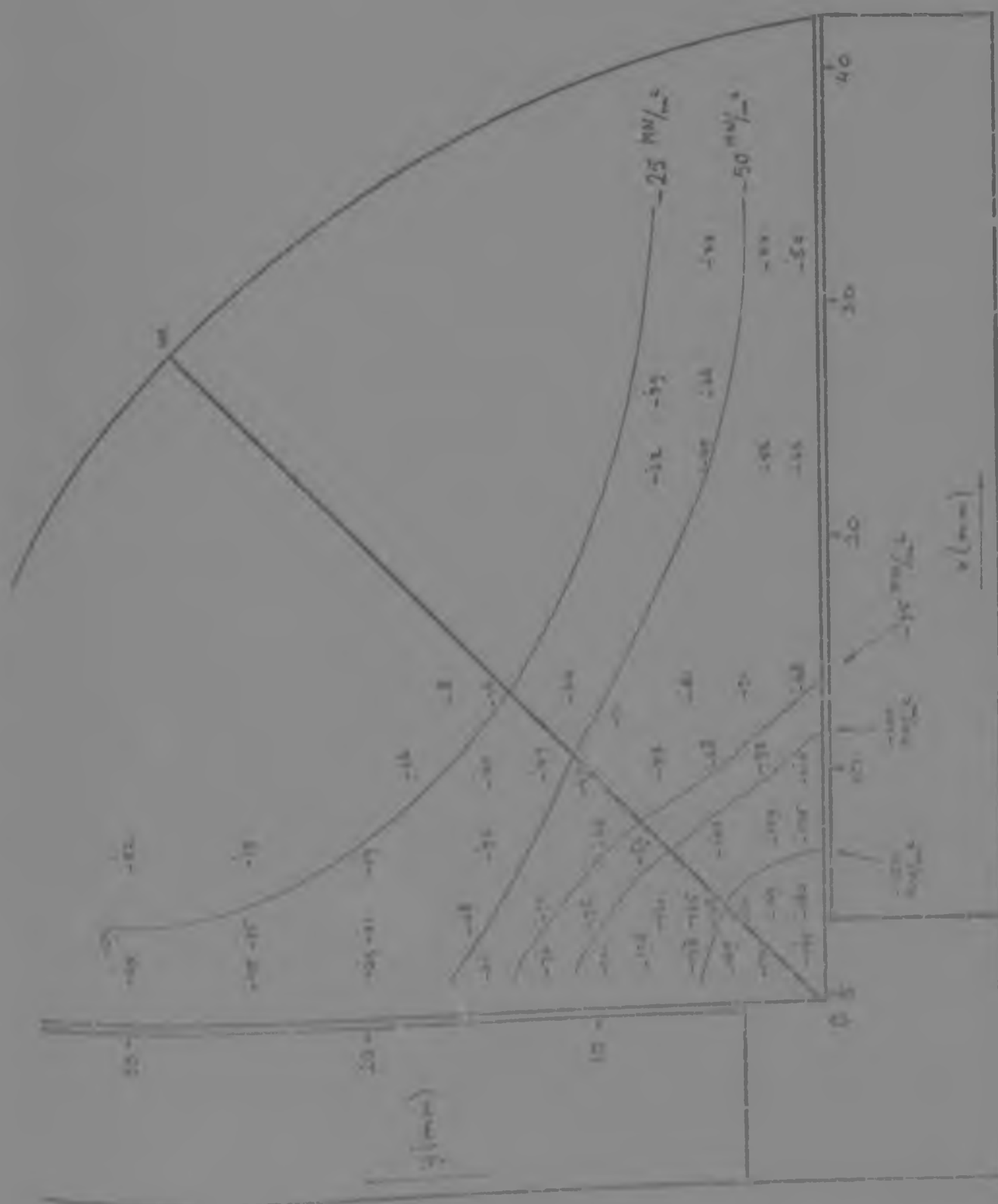


FIGURE 65.

IONIZATION OF THE AIR AT SEA LEVEL BY THE SOLAR RADIATION.
(X-RAY MEASUREMENTS FROM TABLE X).



FIGURE 66.

ISOSTATICS IN THE STEEL SURFACE OF THE ANALOGUE
(X-RAY MEASUREMENTS FROM TABLE XI.)

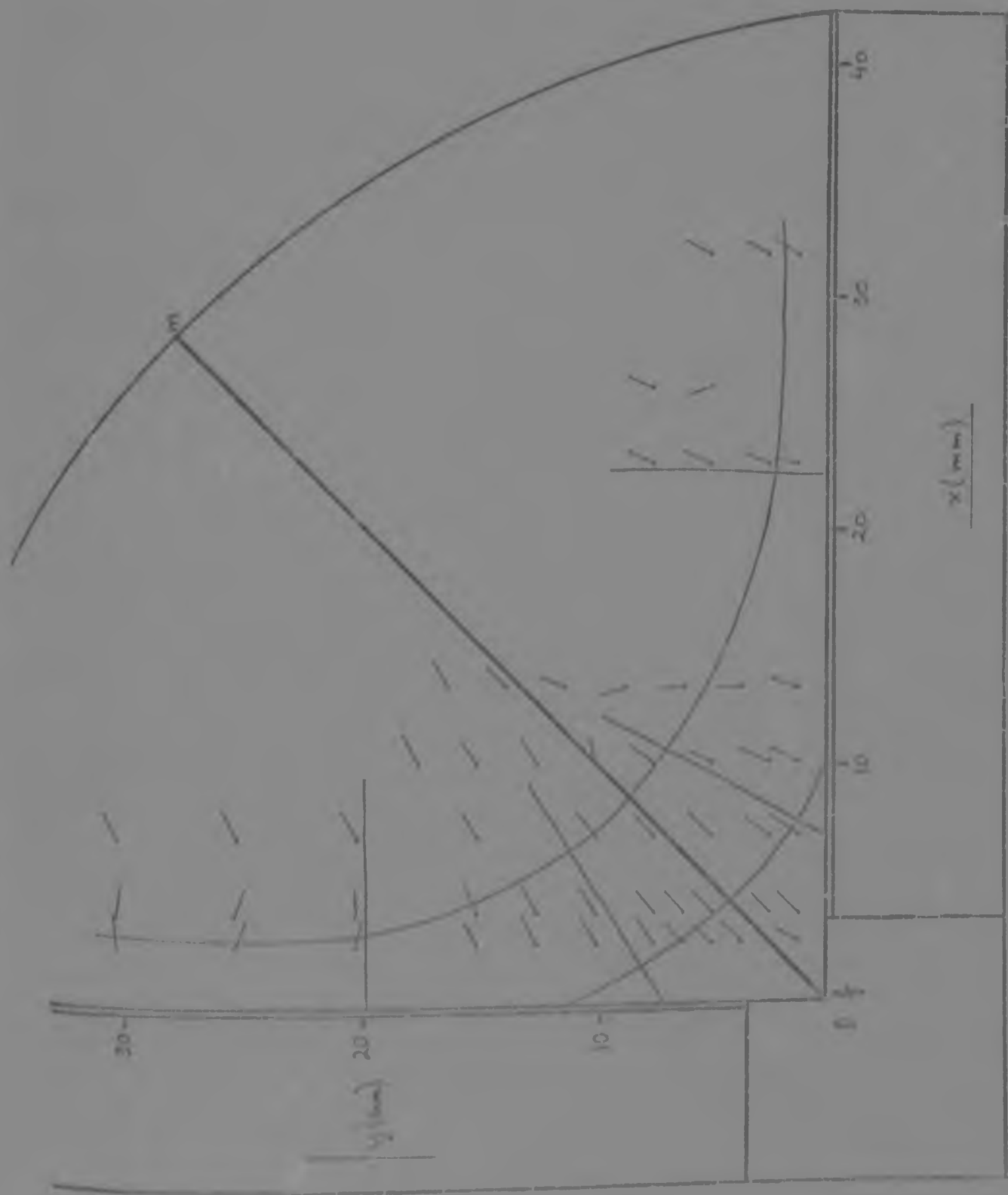


FIGURE 68

FIGURE 70.

ISOCLINICS IN THE STIEL SURFACE OF THE ANALOGUE
(X-RAY MEASUREMENTS FROM TABLE XI).

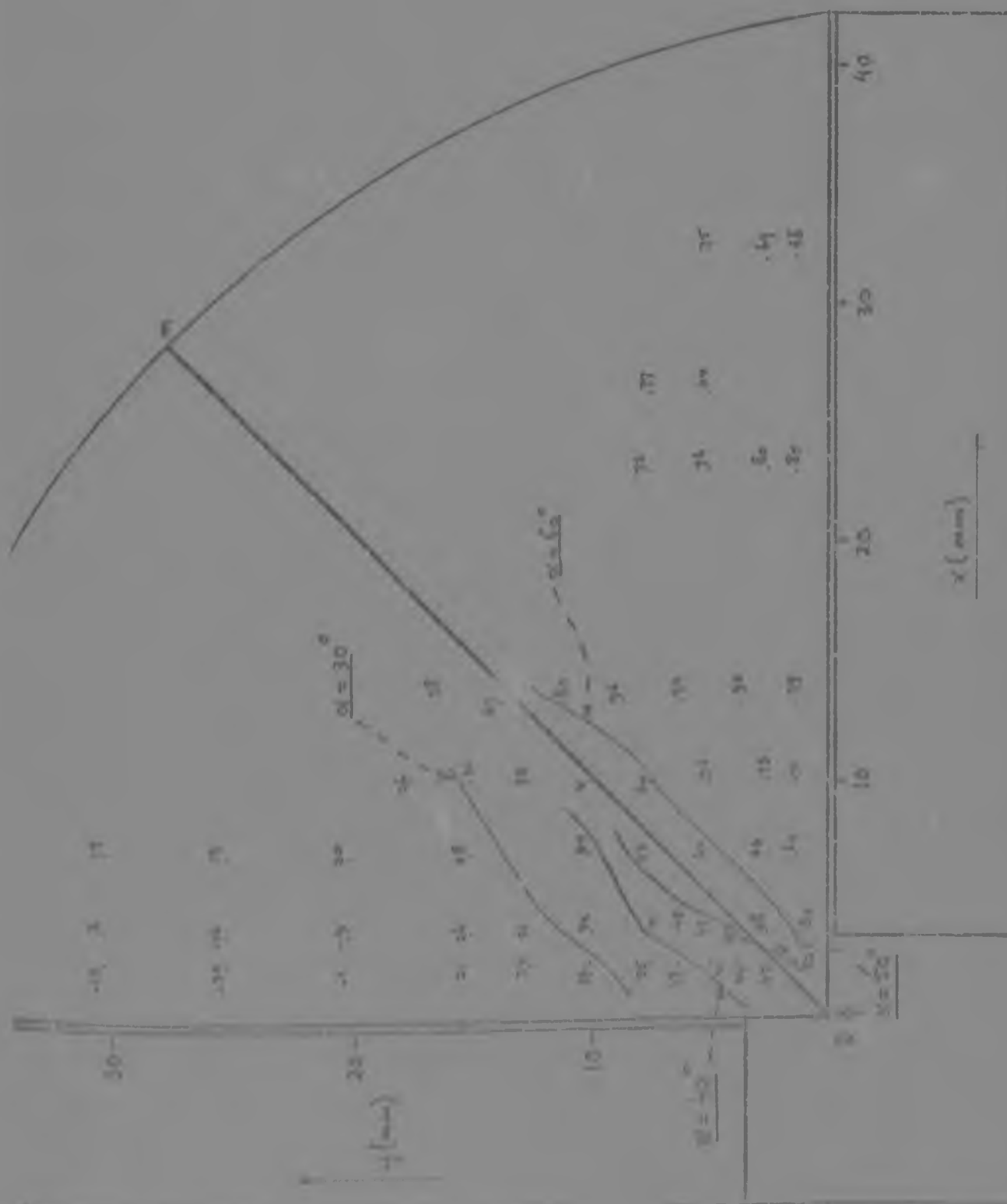


FIGURE 72-

DISTRIBUTION OF PHOTOGRAPHIC ($\sigma_1 - \sigma_2$) ACROSS THE STEEL SURFACE OF THE ANALOGUE, (X-RAY MEASUREMENTS FROM TABLE A1).

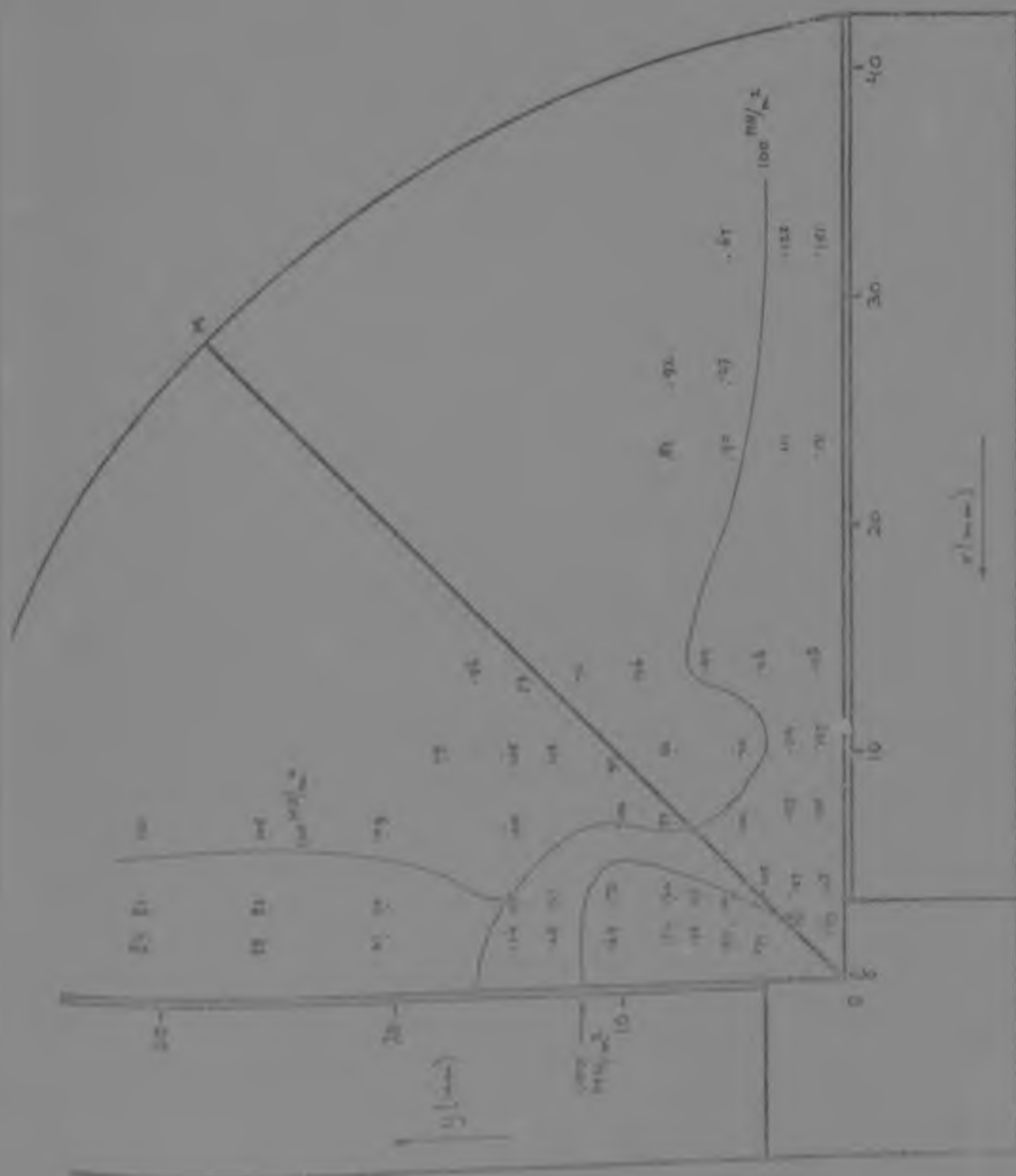


FIGURE 73.

DISTRIBUTION OF MAXIMUM SHEAR STRESS TRAJECTORIES ACROSS THE
STEEL SURFACE OF THE ANALOGUE (DATA FROM TABLE X).

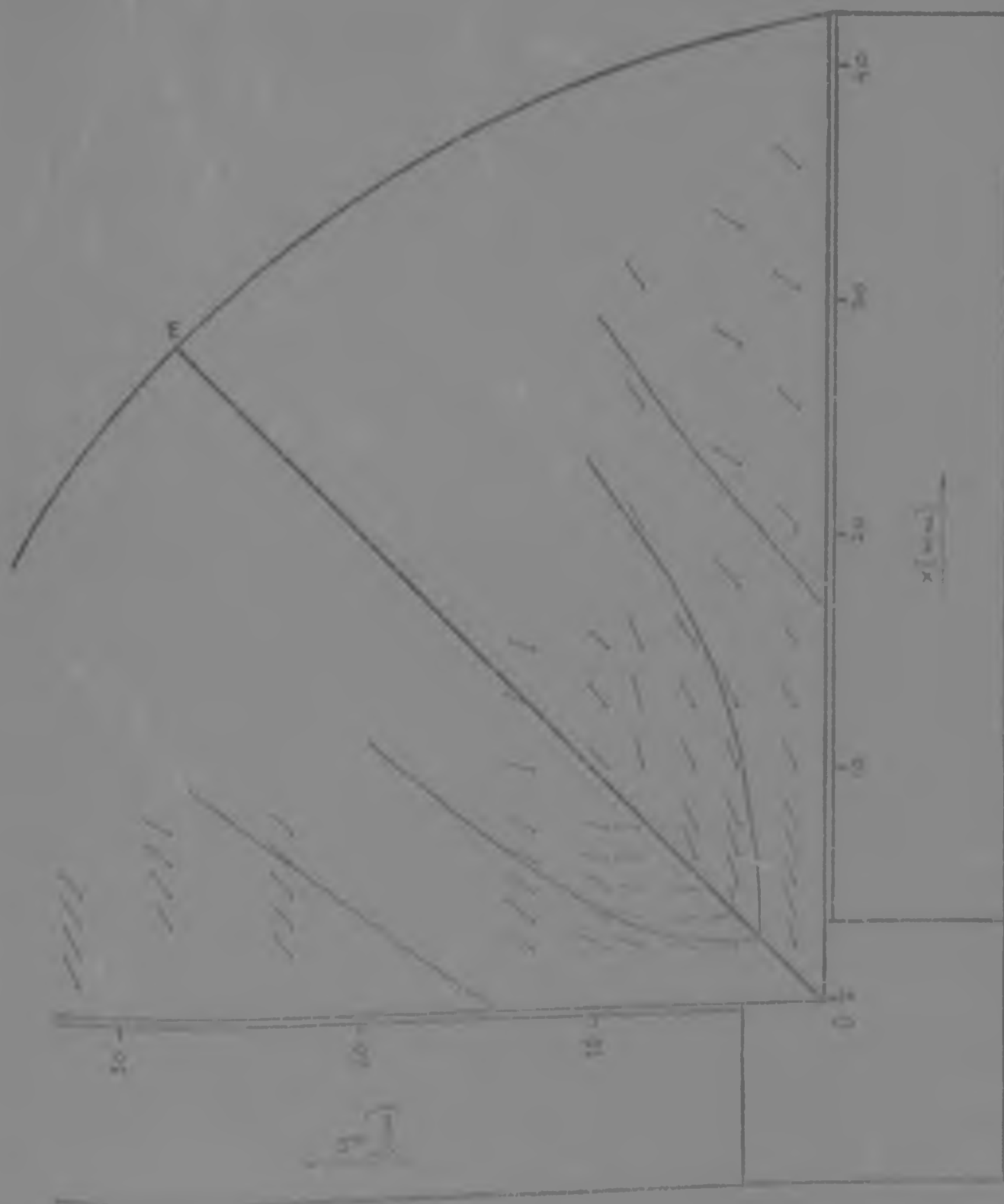


FIGURE 7^a

DISTRIBUTION OF MAXIMUM SHEAR STRESS TRANSFERRED ACROSS THE
STEEL SURFACE OF TOP ANVILS (DATA FROM TABLE XI).

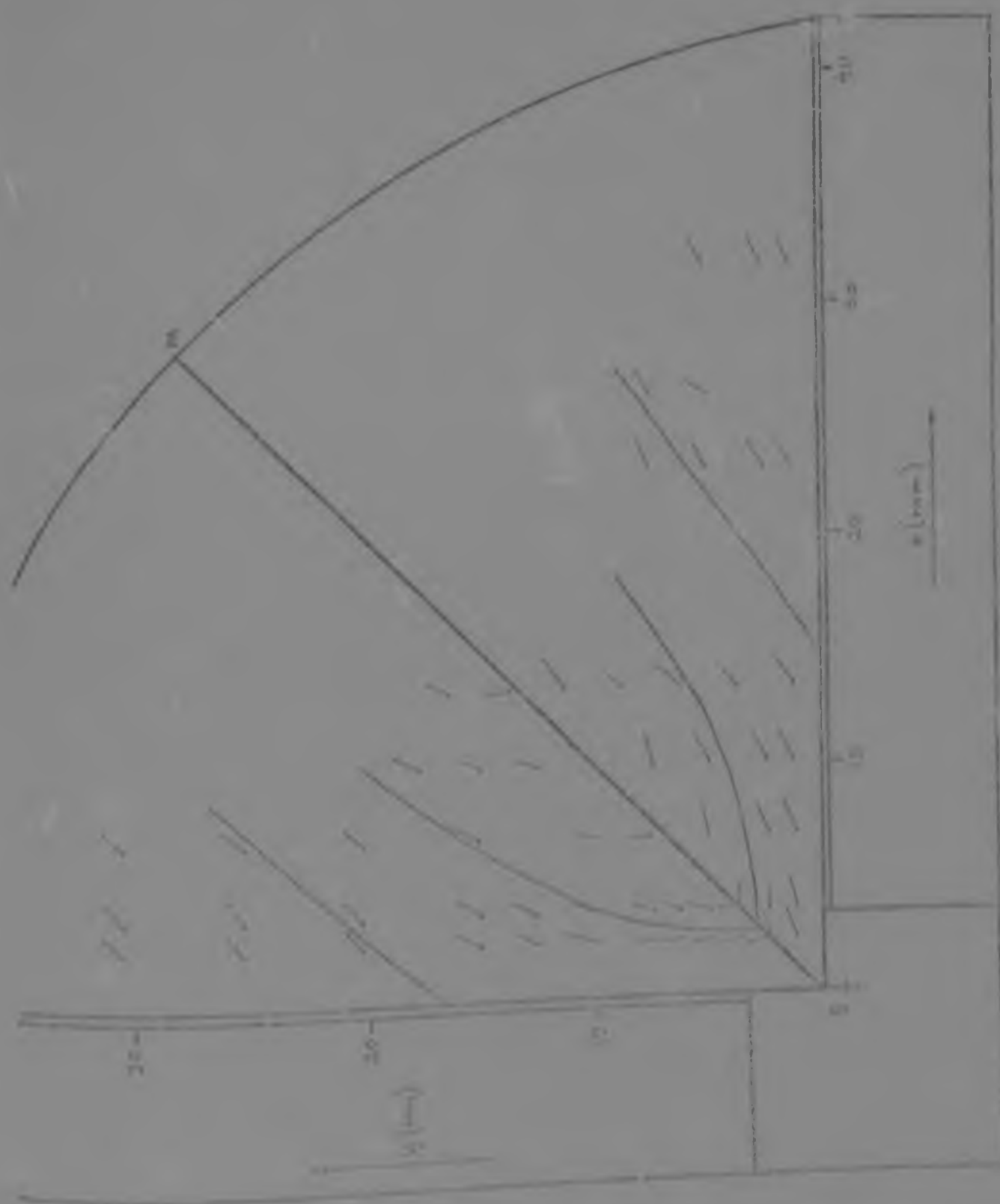


Fig. 75. Finite Element Assembly of a 45°
Sector of the Annular Surface.



FIGURE 76.

RESIDUAL STRESS PATTERN AT 290°C
 DISTRIBUTION OF PRINCIPAL STRESS σ_1 (MN/m^2) ACROSS THE SURFACE
 OF THE ANALOGUE (FINITE ELEMENT ANALYSIS)

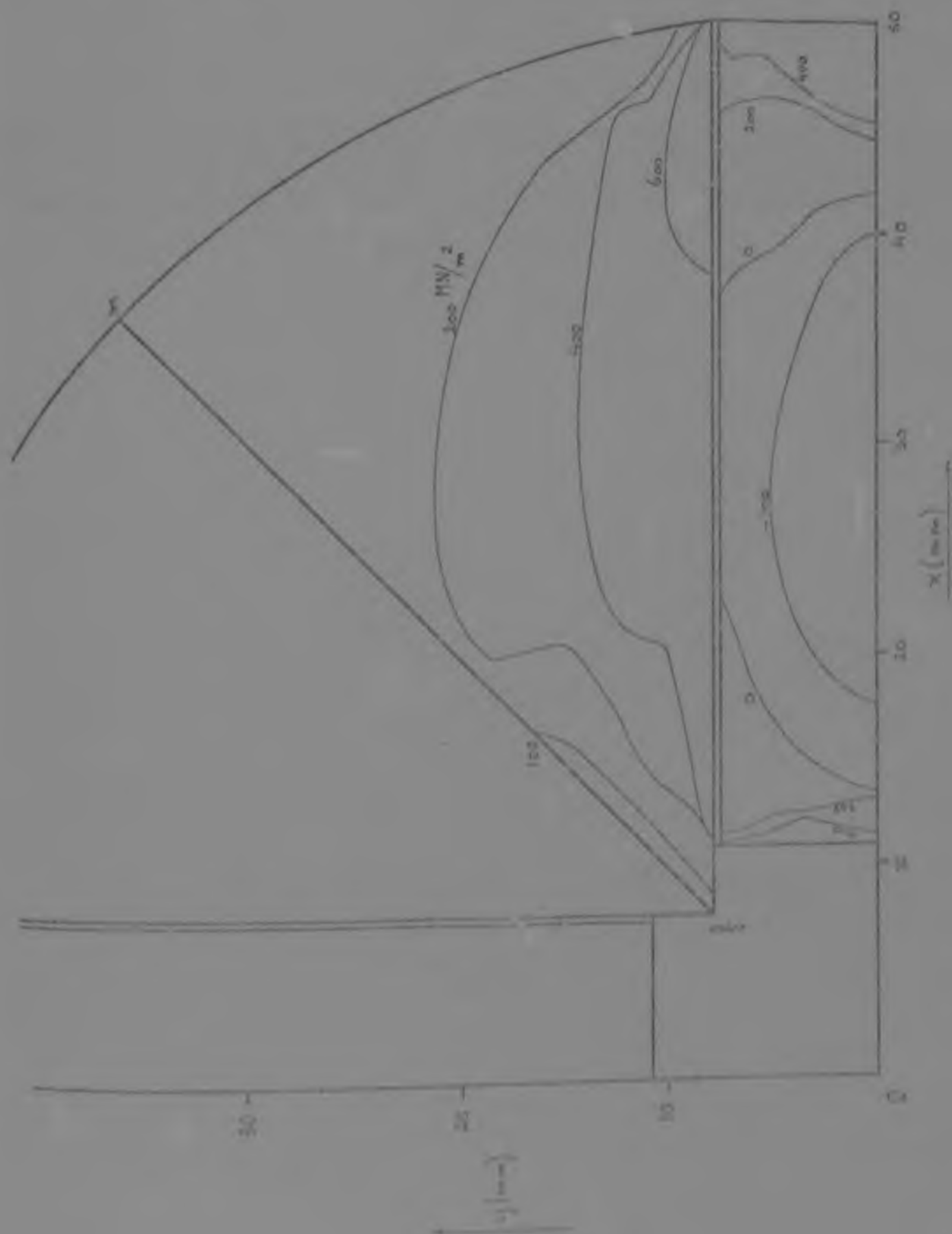


FIGURE 77.

RESIDUAL STRESS PATTERN AT 290°C

DISTRIBUTION OF PRINCIPAL STRESS σ_2 (N/mm²) ACROSS THE SURFACE
OF THE ANALOGUE (FINITE ELEMENT ANALYSIS).

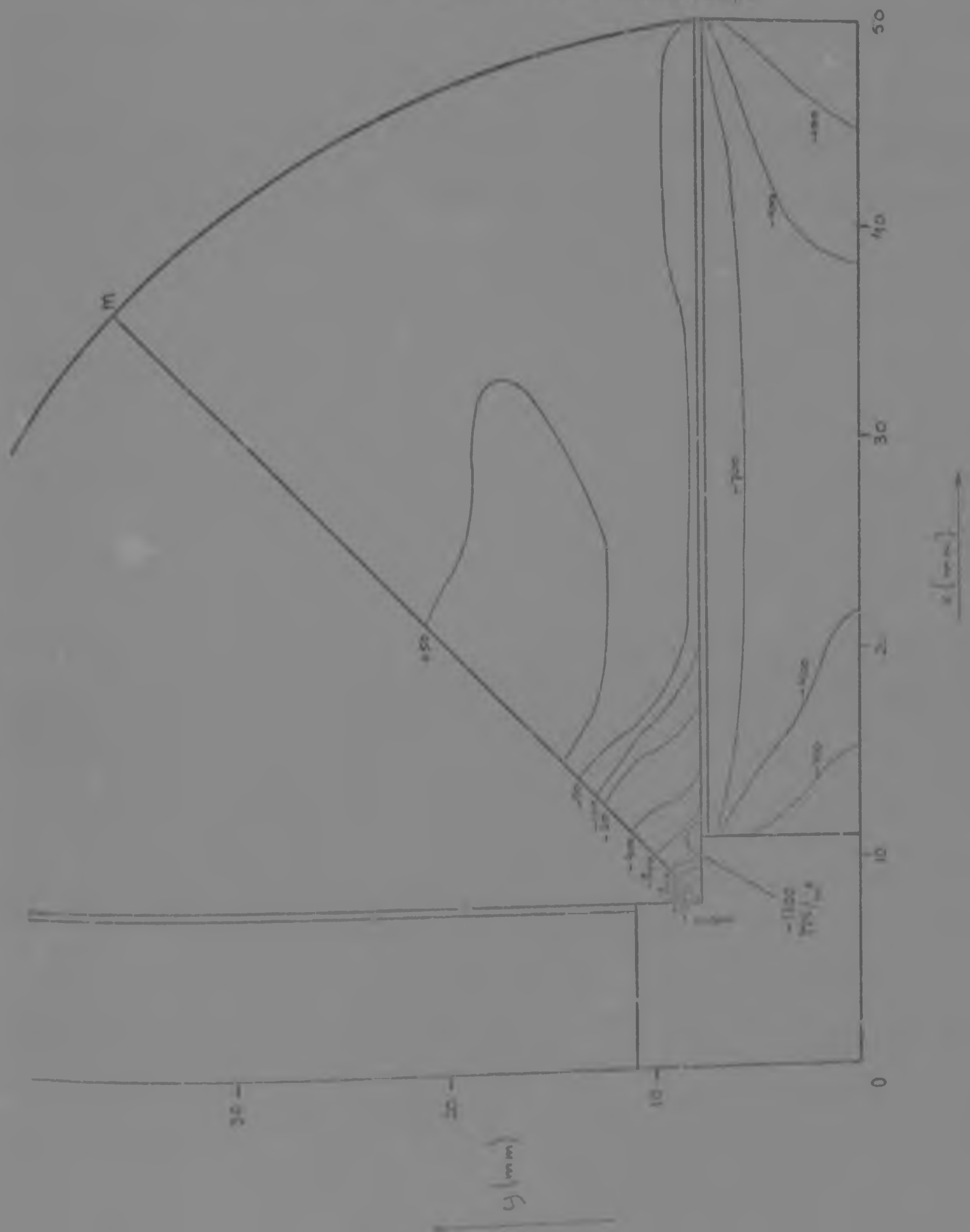


Figure 78.

RESIDUAL STRESS PATTERN AT 170°

DISTRIBUTION OF PRINCIPAL STRESS σ_1 (KSI/IN²) ACROSS THE SURFACE
OF THE AXIALHOLE (FINITE ELEMENT ANALYSIS)

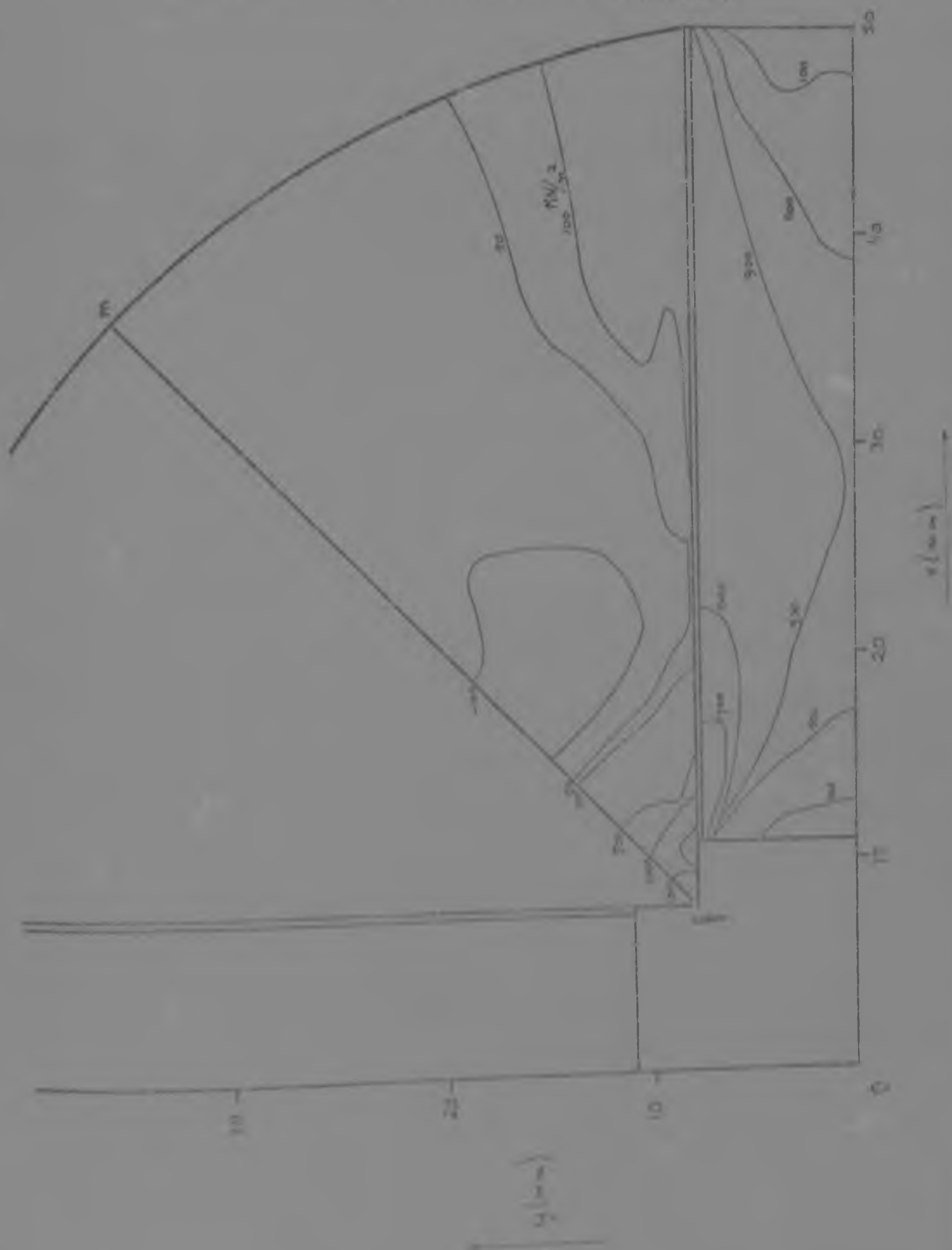


FIGURE 81.

DISTRIBUTION OF PRINCIPAL STRESS σ_2 ACROSS THE SURFACE OF THE ANALOGUE AT ROOM TEMPERATURE. (FROM FINITE ELEMENT ANALYSIS).

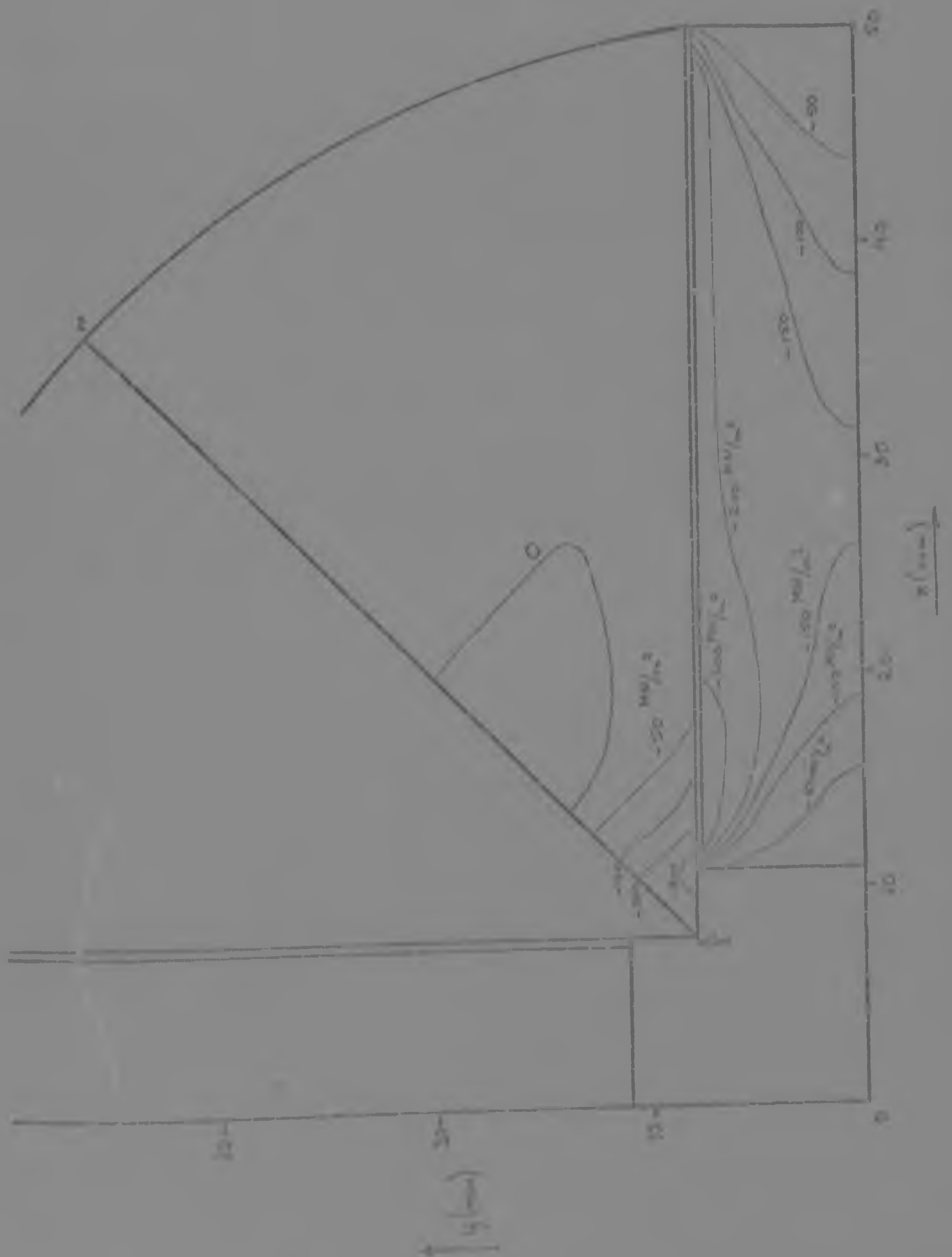


FIGURE 32.

RESIDUAL STRESS PATTERN AT ROOM TEMPERATURE
DISTRIBUTION OF $(\sigma_1 - \sigma_2)$ ISOPHOTOGRAPHIC ACROSS THE STEEL SURFACE
OF THE ANALOGUE (FINITE ELEMENT ANALYSIS).

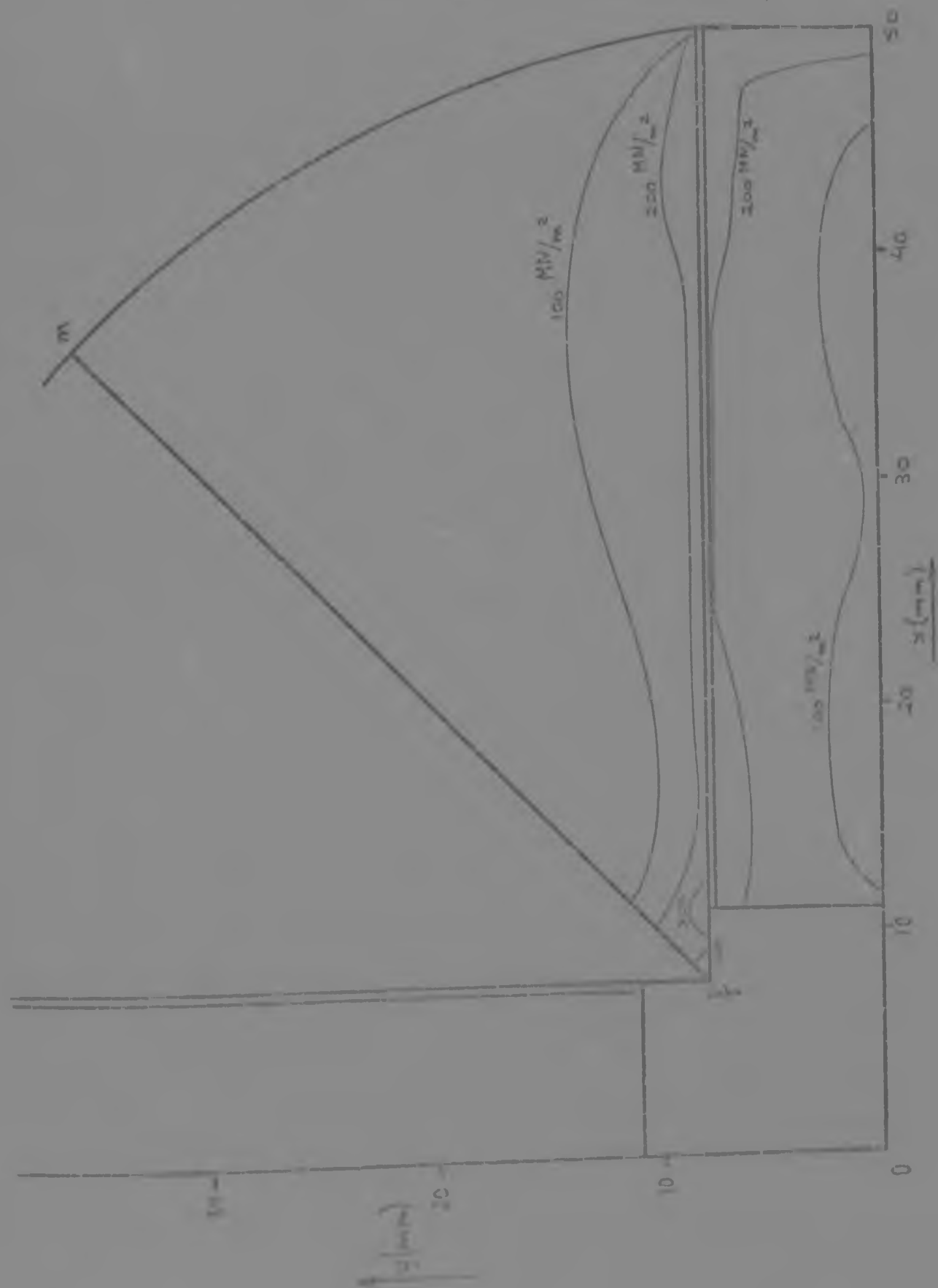
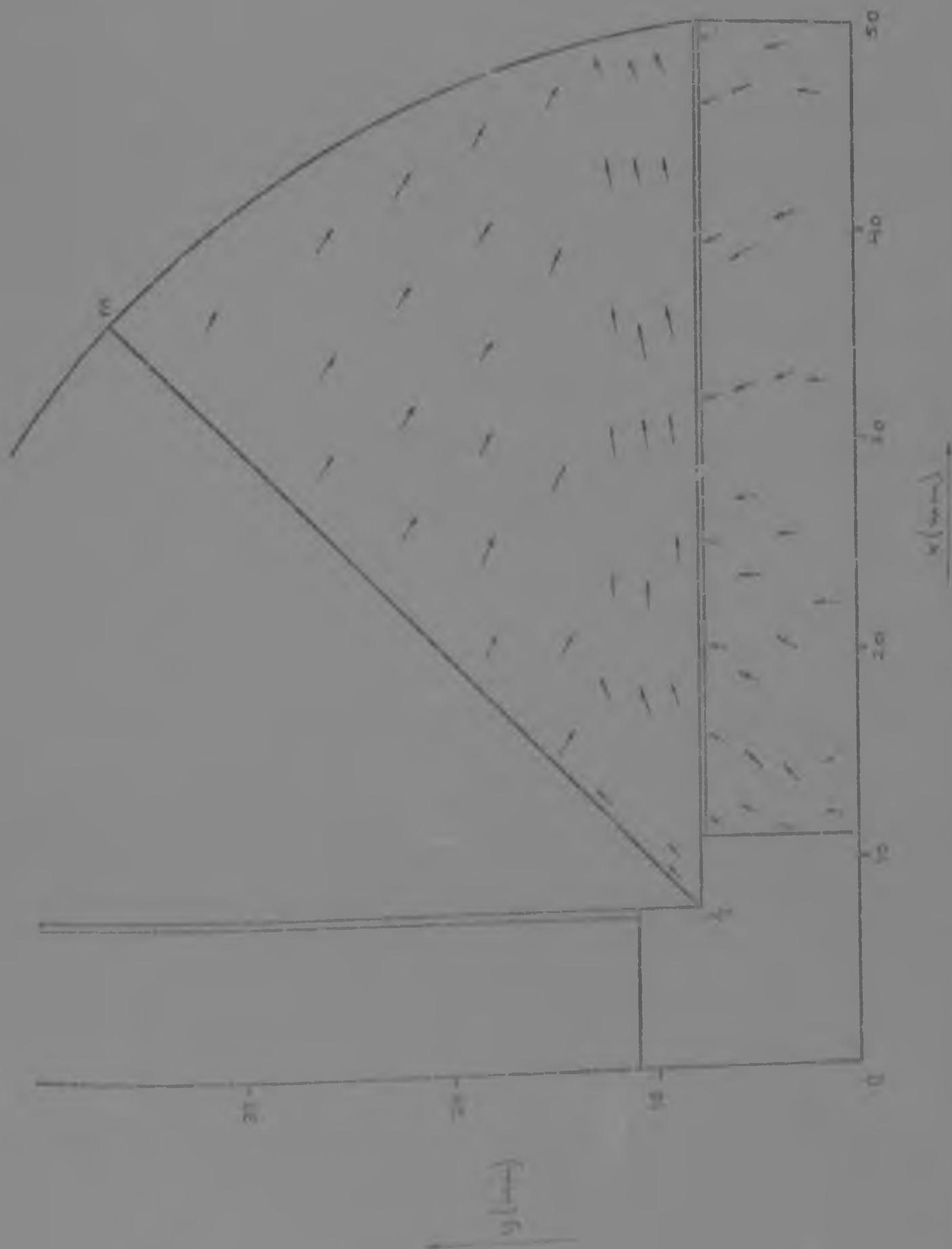


FIGURE 83.

DIRECTIONS IN WHICH THE σ_1 PRINCIPAL STRESS ACTS IN THE SURFACE OF THE ANALOG (AT ROOM TEMPERATURE) (FROM FINITE ELEMENT ANALYSIS).



Author Jonker C G

Name of thesis The application of x-ray diffraction and finite element analysis to the determination of residual stresses and strains on the surfaces of brazed composites of steel and sintered Tungsten carbide-cobalt compacts 1974

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