RADIO-FREQUENCY SIZE EFFECT AND THE FERMI SURFACE
OF ALUMINUM

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Ph. D. Thesis Submitted to Iowa State University

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Radio-frequency size effect and the Fermi surface of aluminum

by

John Sweeney Hartman

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Radio-frequency size effect and the Fermi surface of aluminum

John Sweeny Hartman

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The radio-frequency size effect (RFSE) was used to determine Fermi surface calipers in the (100), (110), and (111) planes for aluminum. The data were recorded using frequency modulation detection and phase sensitive techniques. The measurements were taken on flat, thin (100 - 180 microns) samples prepared from a large single crystal with a residual resistance ratio (RRR) of 13,000. Experimental angular data for each plane were compared with calipers taken from a computer calculation based on the nearly free electron (NFE) model of the second-zone hole surface for aluminum. The calculation yielded relative signal intensities expected for the second-zone signals from purely geometric considerations. These intensities were compared to the experimental results. Comparisons were also made with an adjusted NFE second-zone

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model which included estimates of edge rounding based on previous studies of aluminum.

The effects of non-uniform sample thickness on the frequency study technique for finding the appropriate RFSE magnetic field value were examined. Results of theoretical calculations were compared with data recorded in this investigation. Special consideration was given to the method proposed by Cleveland for determining the frequency dependence of the RFSE line width and the resonant field value. It was found that the Cleveland method gave accurate results only for perfectly flat samples. For samples with non-uniform thickness, the resulting resonant field values were found to be lower than the appropriate value. This reduction was examined as a function of the thickness non-uniformity relative to the skin depth of the material.
I. INTRODUCTION

When undertaking an investigation of the electronic states (Fermi surface) of a metal many experimental techniques are available to the investigator. Methods such as the de Haas-van Alphen effect are well understood and the resulting data can be examined using strong mathematical tools. But the radio-frequency size effect is a relatively new experimental technique and several basic questions remain in interpreting the data.

This investigation was undertaken to use the radio-frequency size effect to examine the well-known Fermi surface of aluminum. By choosing a well-known system the data analysis could also be directed at a better understanding of the radio-frequency size effect itself.

This investigation was aimed specifically at obtaining good measurements on the Fermi surface of aluminum. It was also intended that the aluminum data be used to examine the technique used for selecting the critical parameter from the recorded raw data. This parameter selection has led to uncertainties and discrepancies in past studies and must be evaluated if the radio-frequency size effect is to be utilized to maximum advantage.
II. FERMI SURFACE

A. The Basic Fermi Surface

When the physical properties of a metal are examined it is found that many of them (electrical resistance, specific heat, interaction with electromagnetic waves, etc.) depend at least in part on the behavior of the conduction electrons in the metal. A concise and useful way to describe the states of the conduction electrons in the metal is to introduce the concept of a Fermi surface (FS). Basic concepts behind Fermi surfaces will be discussed and then the FS of aluminum will be examined specifically.

A single electron can be described by a position vector, \( \mathbf{r} \), locating it in the crystal in real space. Similarly it will have a momentum, \( \mathbf{p} = (p_x, p_y, p_z) \), and it can be associated with the point \( (p_x, p_y, p_z) \) in momentum space where the displacement along each axis is proportional to the particle's momentum in that direction. For a classical particle and for free electrons the energy and momentum are related by Equation 1. The mass of the particle is represented by the symbol \( m \).

\[
E = \frac{p^2}{2m}
\]  

(1)

In the real crystal the electrons will occupy the lowest energy states compatible with the Pauli exclusion principle and the temperature of the sample. So for the free electron case where the energy is proportional to the square of the
momentum, the states will be filled beginning at the origin of momentum space. As more electrons are assigned states, they will form spherical shells around the states already filled. For free electrons this leads to the states inside a sphere of radius $p_F$ being occupied and the states outside this sphere being empty. The momentum $p_F$ is called the Fermi momentum. The surface dividing the occupied and unoccupied states is called the Fermi surface and is a constant energy surface. The energy on this surface is denoted by the symbol $E_F$ (the Fermi energy). For free electrons this yields Equation 2.

$$E_F = \frac{p_F^2}{2m}$$

As the temperature of the sample is increased, the separation between empty and occupied states becomes less clear as the probability of occupancy for an electron state near the FS changes. Some states will be occupied above the FS at the expense of some states below the FS. The net effect of this is to cause the FS to become fuzzy rather than being a sharp dividing line between empty and occupied states.

When real metals are examined, the periodicity of the crystal lattice and the interaction of conduction electrons with the fixed nuclei and localized electron core states can greatly distort the free electron model of the FS. The lattice periodicity alone results in the introduction of the concept of a Brillouin zone (BZ) and the associated
modifications in the FS as discussed by Kittel (1) (chapters 3 and 9 respectively). The FSs of most metals actually show little similarity to the free electron spherical FS.

Due to the small mass and resulting small momenta of electrons in a metal, it is convenient to introduce the wavevector, \( \mathbf{k} \), associated with an electron. The wavevector satisfies the following relations

\[ \mathbf{4}\mathbf{k} = \mathbf{p} \quad \text{or} \quad \mathbf{k} = \frac{\mathbf{p}}{\hbar} \tag{3} \]

\[ k = \frac{2\pi}{\lambda} \tag{4} \]

where \( \hbar \) = Planck's constant (h) divided by \( 2\pi \) and \( \lambda \) is the de Broglie wavelength associated with the electron.

Equation 3 indicates the wavevector is proportional to the momentum of the particle. Therefore the FS will be the same shape in \( k \)-space as it is in momentum space. It is standard practice to plot the FS in \( k \)-space. The conversion to \( k \)-space makes it much more convenient to deal with the numbers involved in describing the FS.

Many texts, including Kittel (1, 2) and Ziman (3), discuss in detail the additional factors that determine the actual shape of the FS for a given metal. They also discuss in detail how a knowledge of the FS can be used to calculate physical properties of the material.

B. The Fermi Surface of Aluminum

The Fermi surface of a metal can be studied from two different approaches. Theoretical calculations are carried
out that include all the interactions between particles that the investigators feel are of importance for the metal. But the value of these calculations and their results cannot be evaluated without experimental evidence to substantiate or refute the results of the theoretical models. This exchange between theoretical results and experimental data leads to a better understanding of the physical processes that are actually important for determining the energy states of the conduction electrons in a metal. It also will lead to a better understanding of methods needed to solve difficult problems in the future.

Aluminum forms a face centered cubic (fcc) lattice and the resulting BZ in k-space is shown in Figure 1. Due to the symmetry of the BZ, information on 1/24 of the entire zone is enough for a complete description of the zone and the FS. The portion of the zone chosen is indicated on the figure and several points in the zone are labeled using the notation of Bouchaert et al. (4).

In 1957 Gunnersen (5) used the de Haas-van Alphen (DHVA) effect to study aluminum. The DHVA effect yields information on cross sectional areas of the FS where external (maximum or minimum) cross sectional areas occur (3, p. 274). Gunnersen's data at a given angle were plotted on torque versus (1/H) plots. The curves showed oscillations that were periodic in (1/H). The periods of these oscillations are la-
Figure 1. The Brillouin zone for a face centered cubic crystal structure with the important symmetry points labeled using the notation of Bouchaert et al. (4)
beled $\Delta H$ and are related to the cross section area of the FS.

$$\text{Period} = \Delta(1/H) = \frac{(2\pi e)}{(hcA)} \quad (5)$$

In Equation 5, $\Delta$ represents the extremal area of the FS giving rise to the signal. Heine (6) noted that the angular dependence of the data can be used to deduce the shape of the cross section only if the cross section results from a centrosymmetric piece of the FS. If the cross section belongs to a non-centrosymmetric piece of the FS then we can only deduce a symmetrical surface which has the same angular dependence for the area as the actual FS cross section but the deduced shape need not be correct.

Gunnersen observed signals in two distinct frequency ranges that differed by an order of magnitude. These were referred to as high and low frequency signals. Some authors have referred to them as short period and long period data respectively. Gunnersen associated the high frequency oscillations with pockets of positive holes located at $W$ in the first BZ. This assignment followed his examination of Heine's (6) calculation on the FS of aluminum. Gunnerson's low frequency data were not of sufficient quality or quantity to allow determination of angular dependence of the FS area or an assignment to a specific region of the FS. A study employing several alloys did lead to the conclusion that these data resulted from an electron surface.
Heine (6, 7, 8) carried out an orthogonalized-plane-wave (OPW) calculation and examined the resulting FS in light of Gunnersen's DHVA data and earlier aluminum data taken in studies of the anomalous skin effect (9) and the low-temperature specific heat (10). Heine found the shape of the FS away from the zone boundaries to be very similar to the free electron model. The FS model resulting from this examination had portions of the first three zones occupied. The first zone was completely occupied by electrons except for small regions of holes around the corners of the BZ at W. The second zone consisted of fourteen spherical caps, one on each of the BZ faces. The third zone consisted of small electron surfaces near the corners. Heine's calculation and construction represented the first extensive effort to combine experimental data with energy band calculations to arrive at a FS model.

Fawcett (11) and Langenberg and Moore (12) carried out cyclotron resonance (CR) studies of aluminum and found their results compatible with the theoretical calculations of Heine (6, 7, 8). Two distinct groups of carriers were indicated in each study and Langenberg et al. associated their low mass signals with the holes that Heine had placed at the corners of the first zone.

Harrison (13) then re-examined Gunnersen's data and found that only slight modifications of the nearly free elec-
tron (NFE) FS led to excellent agreement. The calculation began with a construction of the NFE FS by the method that is now known as Harrison's construction. The resulting model is seen in Figure 2. The first zone is full. The second zone contains a single hole surface. The third and fourth zones contain regions of electrons. The NFE model was examined to determine if anticipated DHVA signals would show the symmetry seen in Gunnersen's data. Modifications were made where necessary to arrive at areas compatible with the angular DHVA data. The modifications needed for general agreement occurred in the third and fourth zones. The electron surfaces in the fourth zone were deleted from the model. The shape of the third zone arms was changed but they maintained their original connectivity. The shape Harrison found compatible with the DHVA data is shown in Figure 3a. It was also noted that the sharp edges on the second zone hole surface would be rounded off but no attempt was made to determine how great the rounding would be. The previous interpretation of low mass CR signals would be in error under Harrison's model.

In 1960 Harrison (14) carried out further calculations on the energy bands and the resulting FS using a pseudopotential method. FS cross sections were obtained for several planes passed through the BZ. The cross sections found indicated the rounding expected at the sharp points of the NFE model of the FS due to interactions omitted from the
Figure 2. The NFE Fermi surface of aluminum after Harrison (13) showing the first, second, third, and fourth zones.
Figure 3. Third zone electron arms of the aluminum Fermi surface after a) Harrison (13) and b) Ashcroft (18)
NPE calculation. The cross section shown for a plane containing one of the square faces at $X$ in the BZ indicated the sensitivity of the third zone to small changes in the Fermi energy. Using the Fermi energy found in a single OPW calculation, the third zone model was compatible with the earlier model shown in Figure 3a. But small changes ($\pm 2\%$) in the Fermi energy would cause the arms to swell or be pinched off into small groups of isolated electrons centered at $K$ in the BZ. Due to the sensitivity of the resultant FS at $W$, Harrison could make no definite statement regarding the exact form of the third zone FS based on this calculation. But using the single OPW value of $E_F$ the model was in agreement with areas previously discussed from DHVA data. The model also gave semiquantitative agreement with CR data of Fawcett (11), Moore and Spong (15), and Langenberg and Moore (12) with their low mass data now being attributed to orbits around the third zone arms.

Segal (16) carried out an energy band calculation using the Green's function method. His results were in basic agreement with Harrison's model but he noted that the semiquantitative agreement between the CR effective mass data and the model prediction should not be disturbing. He pointed out that actual masses will depend on electron-electron and electron-phonon interactions which have not been included in calculating the masses from the FS model. It is believed
that these effects would be appreciable and would correct the theoretical value in the direction needed to agree with experimental data.

In 1962 Priestley (17) used the DHVA technique to examine aluminum. An effort was made to find very high frequency signals that would correspond to the second zone hole surface. No attempt was made to reproduce the lower frequency data of Gunnersen (5) and interpreted by Harrison (13). Signals were observed that agreed well with the model of the second zone piece. Also, signals were seen due to electrons moving around the inside of the four arms on a square face. Thus the data gave good quantitative agreement with the NPE third zone model of Harrison.

The main question remaining after this calculation was the connectivity of the third zone arms. Harrison noted that the accuracy of his 1960 calculation was not sufficient to make a definite statement concerning the form of the third zone arms near W.

In 1963 the DHVA data of Gunnersen (5) and Priestley (17) were re-examined by Ashcroft (18) in the development of a new phenomenological model for the FS. The resulting model showed one major difference from the NPE model. Ashcroft proposed that the third zone arms were connected in groups of four around each square face on the BZ but that adjacent groups of arms were not in contact at point W of the BZ. The
proposed third zone model is shown in Figure 3b. Recall that point $W$ is precisely where Harrison (14) could make no definite statement about the FS connectivity due to the extreme sensitivity of that region in his calculation.

In 1967 a definitive experiment for the third zone connectivity was performed by Larson and Gordon (19) using the DHVA method. These data were found to be in excellent agreement with Ashcroft's model.

Additional theoretical and experimental results have been reported but they have not altered the model of the FS appreciably. Many of these works are indicated below but they will not be discussed. Some of these works will be referred to when the data from this present experiment are examined in a later section. Articles marked with an asterisk (*) have already been cited in this work.

Energy band and FS calculations

*Heine (6, 7, 8)
*Harrison (13, 14)
Harrison (20)
*Segal (16, 21)
*Ashcroft (18)
Snow (22)
Greisen (23) (Correction to Snow's result.)
Connolly (24)
Paulkner (25)
Hoffstein and Boudreaux (26)  
de Haas-van Alphen experiments  
*Gunnersen (5)  
*Priestley (17)  
*Larson and Gordon (19)  
Anderson and Lane (27)  
Shepherd et al. (28)  
Cyclotron Resonance experiments  
*Langenberg and Moore (12)  
*Fawcett (11)  
*Moore and Spong (15)  
Galkin et al. (29)  
Naberszhnykh and Tolstoluzhskii (30)  
Spong and Kip (31)  
Magnetoacoustic effect experiments  
Roberts (32)  
Bezuglyi et al. (33)  
Jones (34)  
Fossheim and Olsen (35)  
Kamm and Bohm (36)  
Kohn anomaly experiments  
Stedman and Nilsson (37)  
Weymouth and Stedman (38)  
Galvanomagnetic effect experiments  
Balcombe (39)
Forsvoll and Holwech (40)
Borovik and Volotskaya (41)
Pippard (42)
Balcombe and Parker (43)

Size effect experiments
Brandli et al. (44)
Holwech and Risnes (45)
Risnes (46)

Quantum oscillation experiments
Vol'skii (47)
Balibar et al. (48)
III. THE RADIO-FREQUENCY SIZE EFFECT

A. The Radio-Frequency Size Effect: Concepts

The RFSE is based on the study of anomalies in the surface impedance of single crystal, thin, flat, metal samples in a static magnetic field applied parallel to the surface of the sample. The surface impedance is examined by studying the interaction of the sample with radio-frequency (rf) radiation incident on the sample's surface. Anomalies in the surface impedance are observed for magnetic field values that cause electrons executing cyclotron motion to span the sample thickness and pass within the skin depth on each surface of the sample \(^{(49)}\). For high purity metals at low temperatures, the mean free path of the electrons should be long enough to allow the particle to visit both surfaces before it is scattered. For samples not meeting this requirement, the RFSE is not observed. It will be shown that the magnetic field values corresponding to anomalies in the surface impedance can be related directly to FS dimensions. The RFSE was used in this capacity to examine the FS of aluminum in this investigation.

Throughout the remainder of this work the motion of electrons in real space and in k-space will be examined. Several terms should be explained now to avoid possible confusion in the remainder of the text. The path followed by a particle in real space will be referred to as the parti-
cle's real space trajectory or simply the trajectory. The particle's motion in k-space will be called a k-space orbit or simply an orbit.

As previously stated, the anomalies in the surface impedance are observed when the magnetic field value causes cyclotron trajectories to span the sample thickness. In this situation the electron spends time in the skin depth on both sides of the sample and is able to absorb power from the rf fields in each skin depth. The frequency of the incident rf field is chosen to be much less than the cyclotron frequency. This means that for electrons making more than one cyclotron rotation before scattering, they experience essentially the same field each time they return to the skin depth. Therefore, multiple passes through the skin depth enhance the power absorbed and the electron's effect on the surface impedance anomaly. This effect leads to stronger signals as the mean free paths of the electrons increase. Since the frequencies of the rf field and the electron rotation are not matched, this is not a temporal resonance as in the case of cyclotron resonance. Rather it is the size of the trajectories and the thickness of the sample that are matched so the RFSE is actually caused by a spatial resonance.

The surface impedance anomaly resulting from the cyclotron motion spanning the sample can be related to Fermi surface dimensions by considering an electron's motion in
both k-space and real space. Begin by examining the motion in k-space. The electron is acted on by the standard Lorentz force,

\[ \mathbf{F} = \frac{d\mathbf{p}}{dt} = \mathbf{a} \frac{d\mathbf{k}}{dt} = (e/c)(\mathbf{v} \times \mathbf{H}) = (e/c)(\mathbf{dR}/dt \times \mathbf{H}) \]  

(6)

where \( e \) = charge of the electron, \( c \) = speed of light, \( \mathbf{v} \) = velocity of the electron, \( \mathbf{H} \) = the applied magnetic field, and \( \mathbf{r} \) = position vector locating the electron in the crystal. The force is always perpendicular to the velocity and the magnetic field. The first fact results in the electron's energy remaining constant and the second results in the component of momentum parallel to \( \mathbf{H} \), \( k_H \), remaining constant. In k-space, this restricts the particle's motion to the FS (a constant energy surface) and to a plane perpendicular to the applied magnetic field.

The particle will move around its orbit at the cyclotron frequency,

\[ \omega_c = \frac{eH}{m^*c} \]  

(7)

where \( m^* \) = effective mass of the electron.

Integrating Equation 6 with respect to time leads to Equation 8.

\[ \mathbf{F} \Delta \mathbf{k} = (e/c)(\Delta \mathbf{r} \times \mathbf{H}) \]  

(8)

From this it is seen that the electron's orbit in k-space will be the same shape as the projection of the real space trajectory on a plane perpendicular to the applied magnetic field, but it will be rotated by 90° about the direction of
The projection of the trajectory must be examined because keeping \( k_H \) constant in k-space still allows motion parallel to \( \vec{H} \) in real space. Therefore, in real space the trajectory may be confined to a plane or it may have a helical shape. It is also seen from Equation 8 that for a given value of \( |\Delta \vec{k}| \), the magnitude of \( |\Delta \vec{r}| \) is inversely proportional to the magnitude of the magnetic field. Therefore, as the magnetic field is increased, the projection of the trajectory decreases in size.

In the RFSE, the size limitation placed on the real space orbits is in the direction of the sample normal, \( \vec{n} \), where the sample thickness will be represented as \( t \). In Equation 8 we can then write \( \Delta \vec{r} = t\vec{n} \). With \( \vec{n} \) and \( \vec{H} \) in the previously assigned directions, \( \Delta \vec{k} \) must then be parallel to the sample's surface and perpendicular to \( \vec{H} \). The magnitude of \( |\Delta \vec{k}| \) will then be

\[
|\Delta \vec{k}| = \frac{e}{hc} t H
\]

where \( |\Delta \vec{k}| \) now represents the FS caliper in a direction perpendicular to \( \vec{n} \) and \( \vec{H} \) for the electron orbit under consideration. It is easily seen from this that an accurate determination of the sample thickness and the magnetic field value for the surface impedance anomaly can yield accurate FS calipers. Figure 4 illustrates the relationships between real and k-space orbits and the FS caliper determined by such a measurement. Consider a light beam aimed down the normal
Figure 4. Geometrical relationship between the k-space orbit and real space trajectory for an electron in a magnetic field.
direction, \( \vec{n} \), toward the FS. It is seen that \( |\Delta \vec{k}| \) corresponds to the length of the FS shadow on the line common to the plane perpendicular to \( \vec{n} \) and the plane perpendicular to \( \vec{H} \). For this reason it is possible to take a sample with a \([klm]\) normal axis and use the RFSE to measure the calipers of the FS shadow on the \( (klm) \) plane. As the direction of \( \vec{H} \) is changed in the plane perpendicular to \( \vec{n} \), the calipers for the entire shadow can be measured. It is seen that \( \Delta \vec{k} \) may not correspond to a FS dimension in a plane parallel to the sample surface if the widest points on the FS lie in different planes perpendicular to \( \vec{n} \). This possibility must be kept in mind when data are interpreted.

The use of an rf field incident on the surface of the sample provides a very important contribution by causing the surface impedance anomaly to be relatively narrow. As stated earlier, in order to make a significant contribution to the signal the electron should spend some time in each skin depth. But this can occur for a variety of real space trajectory diameters as illustrated in Figure 5. This illustration is based on a circular electron trajectory to simplify the equations. For a circular orbit, the magnetic field \( \vec{H} \) which leads to a trajectory of diameter \( D \) is given by

\[
\vec{H} = \frac{(2mv\ell)}{(eD)}
\]

where \( v \) = the electron's velocity. For the example in the figure this leads to
Figure 5. Field broadening of the RFSE resonance results from the penetration of rf fields into the skin depth. The real space trajectories shown illustrate the limits of trajectories contributing to the RFSE resonance.
where $H_1$ will be smaller than $H_2$. Consider increasing the magnetic field from a value below $H_1$. When $H_1$ is reached the anomaly in the surface impedance will appear as it is now possible for a complete cyclotron trajectory to occur with the electron visiting both skin depths. As the field is increased further, this trajectory continues to contribute to the anomaly until it is no longer possible for the electrons to enter the skin depth at each surface. This termination of the anomaly occurs at $H_2$. Thus the anomaly is spread over a range of magnetic field $\Delta H$ given by

$$\Delta H = H_2 - H_1.$$  \hspace{1cm} (12)

The fractional width of the signal can be expressed as $(\Delta H / H_1)$ where $H_1$ corresponds to the actual onset of the anomaly.

$$\frac{\Delta H}{H_1} \approx \frac{26}{\pi t}$$  \hspace{1cm} (13)

From this it is easily seen that the anomaly can be sharpened by decreasing the skin depth which can be done by increasing the frequency of the incident rf radiation. For good samples, at low temperatures this can lead to a fractional width of the signals of from one to five per cent.

The idea of sharpening the anomaly by increasing the frequency was first noted by Krylov and Gantmakher (50). It is seen from Equation 11 that the low field onset of the anomaly does not move as the skin depth decreases but all
other features of the anomaly signal will converge toward $H_i$ as $(\Delta H/H_i)$ decreases. This narrowing of the anomaly has been utilized on several occasions to determine the field at the onset of the resonance condition, $H_i$, and is referred to as the frequency study technique.

To carry out a frequency study, RFSE data is taken at several frequencies for the same surface impedance anomaly. In this investigation the data will be plotted on an X-Y recorder where the vertical axis represents the derivative of the surface reactance with respect to the magnetic field and the horizontal axis represents the value of the magnetic field. Figure 6a shows a typical trace from Cleveland's (51) study of molybdenum. The individual features of the signal are labeled. Additional data are taken for the same magnet orientation and range of field values but at several different frequencies on the rf oscillator. The features on the new traces are labeled in the same manner as the original trace and the field values of corresponding features are then plotted on a graph of magnetic field versus frequency to the negative one-third power. The value of the exponent for the frequency follows from the assumption that the skin depth in Equation 13 will be given by the anomalous skin depth. A sample plot is shown in Figure 6b and it is seen that the lines do converge toward a common field value in the limit of extremely large frequency values where the skin depth would
Figure 6. A RPSE data trace (a) and the resulting frequency study (b) are shown from Cleveland's data on molybdenum (51). Numerals correlate RPSE signal features and frequency study lines.
approach zero. This method has been used in several investigations to determine magnetic field values corresponding to the onset of the resonance condition (52, 53, 50, 54, 55, 56).

Throughout this last discussion, the detailed behavior of the electric field in the skin depth has been ignored. The ideas discussed in this semi-classical approach to the problem lead to a good picture of the physical processes involved but do not give information relating to the actual form of the anomaly or the actual rf fields found inside the surface of the sample. Theoretical calculations by Kaner (57) and Juras (58) examine the forms of the rf fields inside the sample and the resulting forms for the surface impedance anomaly.

Another aspect of the RFSE that has been experimentally observed is the coupling of several electron trajectories in the sample to yield a new surface impedance anomaly at a combined magnetic field value. Consider Figure 7 for this discussion. Begin with two FS orbits (#1 and #2) whose anomalies had been observed at field values of $H_1$ and $H_2$ respectively. As the field value is increased both of these trajectories shrink in real space, but some trajectories represented by #1 will still pass through the skin depth at the surface of the sample. The trajectories now pass parallel to the surface again at a depth in the sample given by
REAL SPACE TRAJECTORIES

a. #1
b. #2
c. CHAIN OF #1 AND #2

\[ H_1 \quad H_2 \quad H(#1+#2) \]

Figure 7. The geometry of coupled orbits is shown. Orbits #1 and #2 occur individually of magnetic field values indicated in (a) and (b). The chain occurs at \( H(#1+#2) = H + H \) as seen in (c).
\[ D_i = t \left( \frac{H_i}{H} \right) \] (14)

where \( H = \) current value of the magnetic field. This gives rise to a thin layer of current inside the bulk of the sample which is parallel to the sample's surface (roughly as thick as the skin depth) and contains information about the rf fields at the surface. These current splashes have been found and discussed in the theoretical calculation by Juras (58). This internal current splash can now couple to other trajectories to span the remainder of the sample. For such coupled orbits it is easily shown that

\[ H(\#1+\#2) = H_1 + H_2 \] (15)

where \( H(\#1+\#2) \) is the field value for the anomaly due to the coupled trajectories and \( H_1 \) and \( H_2 \) are the field values for the single trajectories respectively. If a trajectory couples to an identical trajectory, then doubling, tripling, etc. will occur. Takaie (59) observed chains for one orbit containing up to seventeen identical trajectories linked together.

The RFSE was first observed in tin by Gantmakher (49). The evolution of the ideas from cutoff effects in cyclotron resonance experiments to the RFSE is discussed by Takaie (59). Early works using the RFSE were primarily aimed at gaining better FS calipers for a wide range of metals. But as the number of metals left for study decreased, the RFSE technique was turned toward new information. Several good review
articles concerning the RFSE technique have been written (60, 61, 62). RFSE studies have been carried out on the following metals:

- Aluminum (63)
- Antimony (64, 65)
- Bismuth (66, 60, 67)
- Cadmium (68, 69, 70, 71, 72, 73, 74, 75, 76)
- Copper (52, 77, 78)
- Gallium (79, 53, 80, 81, 82, 83, 84, 85, 86, 87)
- Indium (88, 89, 50, 90, 91, 92, 93)
- Magnesium (94)
- Molybdenum (51, 95, 96, 54, 55, 97)
- Potassium (98, 99, 100, 101, 102, 103, 104, 105, 106)
- Rubidium (107)
- Silver (108)
- Thallium (59, 109, 110, 111, 112, 113, 114)
- Tin (115, 116, 117, 118, 56, 119, 49, 120, 121, 122, 123, 124, 125, 126, 127)
- Tungsten (128, 129, 130)
- Zinc (131)

Many of these articles have discussed features of the RFSE mentioned above. Among these are the frequency study technique (72, 52, 53, 50, 54, 55, 98, 103, 113, 114, 56, 119, 108, 57), current splashes or field spikes in the bulk of the sample (86, 107, 58, 57, 132), shapes of the RFSE res-
onance lines (72, 116, 58, 133, 134, 135, 136, 87, 92, 57),
and orbit chains (94, 60, 73, 113, 118, 58, 57).

Recently the RPSE has been turned toward studies other
than PS topology. Among these studies have been
determinations of electron relaxation times (114), skin
depths (82, 50, 87), and electron mean free paths (120, 74,
88, 50, 100, 104, 123, 130, 126, 76, 67, 93, 65, 137, 138,
106). Many of these studies were not aimed solely at these
parameters but gained the information in addition to the
quantities of primary interest to the investigators.

B. The Radio-Frequency Size Effect: Frequency Studies

The concepts involved in the frequency study technique
for determining the correct value of magnetic field for a
given RFSE resonance have been introduced previously. The
technique is based on the narrowing of a RFSE resonance as
the skin depth decreases with increasing frequency. This be-

behavior was first noted by Krylov and Gantmakher (50). Since
then it has been used by many (72, 52, 53, 50, 54, 55, 98,
103, 113, 114, 56, 119, 108, 57) in an effort to determine
the correct magnetic field values to use in Equation 9.

Several recent RFSE studies have been performed on
molybdenum (95, 96, 54, 55, 51, 139) with differing results.
The PS calipers obtained in these studies show the same
angular dependence except they differ in magnitude by about
five per cent. A recent DHVA study of molybdenum (140) has
found good agreement with Boiko et al. (95, 55, 139). Boiko et al. in their early work chose the magnetic field value corresponding to the low field end of the resonance feature. Cleveland (51) and Cleveland and Stanford (96, 54) made extensive use of the frequency study technique to determine field values for the different resonances observed. The use of the frequency studies apparently led to values of the magnetic field that are five per cent too small. Figure 8 shows a Cleveland data trace (51, Figure 35) and includes two arrows marking the values of magnetic field for the resonances as found in the frequency study technique ($H_{res}$) and also arrows indicating the first departure of the RPSE signal from the background signal ($H'$). Notice that these resonance field values fall at lower field values than the first departure of the RPSE signals from the background signal. Based on the results of these molybdenum studies it would be useful to re-examine the way in which the frequency studies were carried out by Cleveland.

RPSE data were taken with all experimental conditions the same except the frequency of the rf oscillator. From these data traces, individual features were labeled as in Figure 6a. Then the field value at each feature and the uncertainty in that field value were determined by measuring the trace relative to the known end points on the field sweep. The values of magnetic field ($H$), uncertainty in
RF SIZE EFFECT IN Mo
SAMPLE NORMAL = [110]
SAMPLE THICKNESS = 5.10 x 10^{-3} INCH
f = 10.05 MHz
H \perp [100] AXIS

Figure 8. Cleveland data trace for Mo, showing the resonance magnetic fields found from frequency studies and the magnetic fields where the signal first deviates from the background curve.
field ($\Delta H$), and frequency ($f$) were then known for each feature measured. The information for the same feature at all different frequencies was grouped together as these points will contribute to a single line on the frequency study as seen in Figure 6b.

The data for each line were then used to carry out a linear least-squares fit to a line given by

$$H = A + B f^{-\alpha}$$

(16)

where $A =$ intercept of the line at $f=\infty$, $B =$ slope of the line, and $(-\alpha) =$ the exponent used for the frequency. For this linear least-square fit each point was given a weight inversely proportional to the square of the uncertainty in the field value for the point (141). This process led to values of $A$ and the uncertainty in $A$ for each line at a given value of $(-\alpha)$. For all lines in a group (such as Figure 6b) the values of the intercepts were statistically combined to yield an average intercept $A_{\text{ave}}$ and the uncertainty in that intercept $\sigma_A$.

This procedure was repeated as $\alpha$ was incremented over a range of values that included the frequency exponent for the classical skin depth ($1/2$) and the anomalous skin depth ($1/3$) for a free electron gas. The results were plotted on graphs of $\sigma_A$ versus $\alpha$. The value of $\sigma_A$ was taken to be the size of the error bar for the value of the average intercept with a 90% confidence level. For the frequency study on the six
lines of Figure 9a (Cleveland Figure 30) the resulting plot of $\sigma_A$ versus $\alpha$, Figure (9b), showed a minimum at $\alpha=0.33$. This was taken as confirmation of the agreement with the anomalous skin depth frequency dependence. The values of $A(\text{ave})$ for the remaining frequency studies were then determined for $\alpha=0.33$.

Several points should be mentioned concerning these linear least-squares fit calculations.

1) The values of chi-squared (141) for the individual lines are very insensitive to changes in $\alpha$ for the data taken. The chi-squared values were often smaller than unity for large ranges of $\alpha$ and therefore were not a meaningful indicator statistically for choosing a preferred value of $\alpha$. This results from a relatively small range of frequency data, a long distance of extrapolation, and relatively large error bars on the values of magnetic field. Based simply on the value of chi-squared it would be impossible to choose a preferred value of $\alpha$ to give a "best" fit for a given line.

2) The values of $\alpha$ corresponding to minima in the $\sigma_A$ versus $\alpha$ curves do not all have the same magnitude. They are spread over the range from 0.25 to 0.44. The wide range of values is not encouraging and some of these values (from 0.33 to 0.25) fall outside the
Figure 9. Frequency study (a) and the resulting confidence limits for the mean value of the intercept as a function of the exponent for the frequency dependence of the RFSE line broadening (b). Original data after Cleveland.
range determined by the classical and anomalous skin depths. The groups of lines examined with three or more lines per group exhibited a smaller range of $\alpha$, from 0.32 to 0.35.

3) Even if all values of $A(\text{ave})$ were chosen for $\alpha=0.33$ the field corrections would not be large enough to account for the five per cent difference with the other molybdenum studies. In most cases the change would be less than one per cent.

From this brief examination it is evident that the concept of the frequency study and Cleveland's method for determining the proper frequency dependence should be examined more closely. One experimental parameter that has not been examined in relation to the frequency study is the uniformity of the sample's thickness.

Consider a wedge-shaped sample whose thickness varies between $t$ and $t+A$. Examine motion for an electron making a circular trajectory in the sample which is subjected to an rf field at the surfaces. A cross sectional view of the sample and the possible orbits is shown in Figure 10. As was seen in the earlier discussion of RFSE signal broadening due to the skin depth, it is again possible for a variety of trajectory diameters to expose the electron to the rf fields in the skin depth at both surfaces without striking either surface. The two orbits illustrated represent the maximum
Figure 10. The effects of variations in sample thickness and the skin depth on the variety of electron trajectories that can contribute to the RFSE are illustrated.
and minimum diameter orbits that can do this and they satisfy the conditions

\[ 2r_1 = (t + \Delta) \quad \text{and} \quad 2r_2 = (t - 2\delta). \quad (17) \]

Following steps analogous to those used in the earlier discussion of line broadening, this leads to the conclusion that this electron orbit can contribute to a RFSE resonance over a range of magnetic fields. The fractional width of the RFSE resonance can be written as

\[ \Delta H/H_1 = \left( \Delta + 2\delta \right)/t. \quad (18) \]

The effect of this broadened contribution to the RFSE resonance must be examined to determine the effect expected on the signals observed experimentally.

For the electron trajectory considered, the electron will be able to interact with the rf fields and contribute to surface currents and surface impedance equally well for all values of magnetic field between \( H_1 \) and \( H_2 \). The resulting effect on the surface impedance is schematically represented in Figure 11a. But the RFSE signals are proportional to the field derivative of the surface impedance and may be expected to show some similarity to the curve shown in Figure 11b. Notice the splitting into two signals and the inversion of the two signals on the derivative plot relative to each other. This is exactly the kind of behavior mentioned by Gantmakher (60, Figure 19) for a wedge-shaped sample. This simplified, classical argument has again neglected the actual
Figure 11. A schematic representation of the effect of a wedge-shaped sample on the surface impedance is shown (a). The resulting RFSE signal detected in this investigation is shown (b).
form of the rf fields in the sample and should be used only to gain insight into the type of behavior expected. Exact agreement with experimental line shape should not be expected.

Several authors (60, 74, 50, 119, 108, 135) have noted that a non-uniform sample thickness can lead to signal broadening rather than signal splitting as seen above. Consider the case where the sample surfaces are undulated rather than simply flat but non-parallel. Then for different values of magnetic field between $H_1$ and $H_2$ different numbers of electron trajectories may contribute to the surface currents and surface impedance. One possible form of the resulting surface impedance is shown in Figure 12a and the resulting field derivative is shown in Figure 12b. The exact forms of these curves would depend on the actual pattern of thickness variations in the sample.

These two cases have indicated that non-uniform sample thickness will generally lead to broadening of the RFSE signal and distortion of the line shape from the form that would be observed in a perfectly flat sample. For the special case of a perfectly wedge-shaped sample, RFSE signal splitting will occur.

From Equation 18 it is seen that the non-uniformity in sample thickness and the skin depth affect the RFSE resonance width in the same functional manner. The relative magnitude
Figure 12. a) A schematic representation of the effect of random variations in the sample thickness on the surface impedance is shown. b) The resulting RFSE signal detected in this investigation is shown.
of the two factors can be evaluated by examining a frequency study plot and Equation 18. For infinite frequency, the RPSE line width is governed by the non-uniformity of the sample thickness.

\[ \frac{(\Delta H)}{H} = \frac{\Delta}{t} \]  

(19)

Then at a finite frequency, the line width is the sum of the frequency independent term given in Equation 19 and the standard broadening due to the skin depth as first seen in Equation 13. If the skin depth is taken to be

\[ \delta = a|f - \alpha| \]  

(20)

then the skin depth broadening is directly proportional to the coefficient "a" in Equation 20. Thus the slope of the steepest line on a frequency study is directly proportional to the coefficient seen in this simplified expression for the skin depth.

The line width at any frequency can be broken into the width due to the skin depth and the width due to thickness variations. The line width due to the skin depth is proportional to the difference in the slope of the steepest and flattest lines in a given frequency study. For example in Figure 9a, the slopes would be measured for lines I and VI. The difference in these slopes is then an indication of the importance of the skin depth in broadening the RPSE signal. The line width due to sample thickness variations is found by examining the frequency study lines for infinite frequency.
The spread in their intercepts is an indication of the importance of thickness variations in broadening the RFSE signal. The division of the largest difference in the intercepts by the largest difference in slopes results in a number that indicates the relative importance of the thickness broadening and skin depth broadening. The quantity resulting from this division has been called RATIO and will be used on several figures. As RATIO decreases the thickness broadening becomes less important relative to the broadening due to the skin depth. RATIO is equal to zero for a perfectly flat sample and the skin depth is the only source of line broadening. As RATIO increases, the sample quality is decreasing as the sample behaves less and less like the thin, flat sample wanted ideally.

The effect of this broadening on Cleveland’s method for determining the proper frequency dependence for the line width and the correct field value has been examined. This was done by generating data points that fell on lines showing the frequency dependence seen in Equation 16. Systematic variations were included in the intercepts of the individual lines and in their slopes. These variations thus incorporated changes in the flatness of the sample and the magnitude of the rf skin depth. Data were generated for five frequencies between 2.4 MHz and 18.6 MHz.
The generated data sets were examined using the Cleveland linear least-square fit method for the individual lines. The value of the frequency exponent, $\alpha_{\text{min}}$, was found that gave the least uncertainty, $\sigma_\alpha$, in the common intercept, $A_{\text{ave}}$. This corresponds to the $\alpha$ at the minimum on a $\sigma_\alpha$ versus $\alpha$ plot as seen in Figure 9b. Figure 13 illustrates the dependence of $\alpha_{\text{min}}/\alpha_{\text{real}}$ on RATIO, the relative importance of the thickness broadening compared to the skin depth broadening. Data are shown for $\alpha_{\text{real}}=1/3$ and $1/2$. Both data sets show a very similar dependence on RATIO with only small differences in the value of $\alpha_{\text{min}}/\alpha_{\text{real}}$ for a given value of RATIO. From this it is seen that for perfectly flat samples, RATIO=0, Cleveland's technique yields the proper value of the exponent seen in Equation 16. The value of $\alpha$ indicated by the Cleveland method decreases as the sample becomes less flat. Any variation in sample thickness will cause the resulting $\alpha$ to be lower than the actual value.

This calculation showed that $\alpha_{\text{min}}/\alpha_{\text{real}}$ depended only on the relative importance of the two sources of broadening and not their absolute magnitude.

The effect of a non-uniform sample thickness on the common intercept resulting from the Cleveland technique, $A_{\text{ave}}$, has also been examined. The common intercept is examined relative to the lowest intercept of the frequency
Figure 13. The theoretical dependence of \( \frac{\alpha_{\text{min}}}{\alpha_{\text{real}}} \) on the parameter RATIO in the Cleveland (51) frequency study technique is illustrated.
study lines when they are examined using Equation 16 with the frequency exponent taken as \( \alpha (\text{real}) \). This lowest real intercept will be denoted \( A(\text{min}) \). Figure 14 illustrates the dependence of \( (A(\text{min})-A(\text{ave})) \) versus the parameter RATIO. Positive values on this vertical scale represent amounts that \( A(\text{ave}) \) is reduced below \( A(\text{min}) \). From the figure it is seen that the reduction depends only on spread in the intercepts of the lines in the frequency study. It can also be noted from Figure 14 that the value of \( (A(\text{min})-A(\text{ave})) \) is roughly the same size as the spread between the intersections for the lines in the frequency study. The points are grouped about horizontal lines so they show no functional dependence on RATIO. Hence the reductions do not depend on the skin depth or its frequency dependence.

This behavior can easily be understood by considering the form of the data. First consider the data plotted on a graph of magnetic field versus \( f^{-\alpha} \) where the actual exponent for the data is used. Assume the case where two frequency study lines exist and they do not have a common intersection. If the value of \( \alpha \) for the plotted lines is increased without changing the horizontal scale, the lines will bend upward in the region near the vertical axis. The upper line will bend up more than the lower line because its slope will be larger than that for the lower line. Thus the intersections will diverge as the exponent is increased. This is a direct
Figure 14. The dependence of the reduction in the average frequency study intercept on the actual spread in line intercepts and the parameter RATIO.
result of the fact that for the data being considered, $f^{-\alpha}$ is always less than unity.

Next consider decreasing the value of $\alpha$ for the plotted lines. Again leave the horizontal axis unchanged. Now the two curves bend downward in the region near the vertical axis. The upper line is again bent more because of its larger slope. Thus the intersections for the two curves converge and for some $\alpha(\text{min}) < \alpha(\text{real})$ the spread between the intersections will be minimized. This corresponds to the value of $\alpha(\text{min})$ selected by the Cleveland technique. It is seen that any spread between the intersections will cause the indicated $\alpha(\text{min})$ to be less than $\alpha(\text{real})$. It is also seen that the common intercept at $\alpha(\text{min})$ will be smaller than the actual intercept found for $\alpha(\text{real})$.

The effect of a non-uniform thickness has been examined on both the exponent and the intercept resulting from Cleveland's frequency study method. Deviations in sample thickness have been found to cause the resulting values of $\alpha(\text{min})$ and $A(\text{ave})$ to be lower than the correct values.

Throughout this discussion the rf field has been assumed to have only a single frequency component. The affects of possible frequency harmonics present in the rf fields have been ignored.

The presence of harmonics in the rf field would further complicate the frequency study technique. Suppose the rf
fields at a sample's surface have components at $f_o$ and $3f_o$. By examining a frequency study plot it is seen that the $3f_o$ RFSE resonance will occur at a lower magnetic field value than the $f_o$ resonance. The two resonances would combine in some manner. The resulting resonance could be expected to begin at the low field end of the $3f_o$ resonance and extend to the high field end of the $f_o$ resonance as seen in Figure 15, but the frequency counter would indicate $f_o$. The result of this would be a frequency study plot where the data points would be located at magnetic field values that would be artificially low. The confusion could be compounded if the harmonic content of the rf signal depended on the oscillator frequency as seen for tube oscillator A. In this case the amount of shift toward lower field values would be proportional to the amount of harmonic content in the rf signal.

From the preceding discussions, several conclusions can be reached about the frequency study technique for determining the magnetic field where the RFSE resonance begins.

1) Efforts must be made to ensure that the rf signal applied to the sample is a pure sine wave with virtually no harmonic content.

2) Variations in the sample's thickness can lead to signal broadening and a finite RFSE line width for infinite frequency. To avoid this, efforts must be directed at minimizing the deviation of the sample from
Figure 15. The effect on the surface impedance is illustrated for a RFSE resonance exposed to a signal containing frequencies $f_0$ and $3f_0$. 
a flat plate. Using small area samples may be helpful for this. If the two surfaces of a sample are slightly misaligned, the variation in thickness is then proportional to the length of the flat surfaces. A 0.5° misalignment between the two surfaces can result in a difference of 70 microns in the thickness at opposite ends of an 8 mm sample. Therefore smaller sample area leads to smaller variations in the sample thickness.

3) Efforts must be directed toward minimizing the uncertainty in the field values of individual line features. This would allow a better determination of the frequency dependence of the individual lines in a frequency study group. Smaller error bars on the magnetic field would make the chi-squared values more useful for the individual lines. This could be done by taking duplicate data traces and determining the average magnetic field for a feature and its standard deviation.

4) Extending the frequency range studied would also help make the chi-square values more useful for determining the frequency dependence of the individual lines.
IV. RADIO-FREQUENCY SIZE EFFECT SIGNALS IN ALUMINUM:

COMPUTER PREDICTIONS

A serious problem can exist in RFSE experiments when one attempts to associate individual signals with the appropriate Fermi surface calipers. Roach (94) emphasized this in his discussion of the RFSE technique and the FS of magnesium. Many signals were seen and orbit assignments would have been very difficult without an accurate FS model. This problem is compounded by the fact that orbits can form chains to give signals at magnetic field values that do not correspond to a single FS caliper. This effect can give rise to doubling (or tripling, etc.) if the orbits are the same or to totally new shapes on the plot of caliper versus magnet angle if two different orbits couple to yield one signal. Roach even noted an orbit that did not give rise to a signal by itself but did appear coupled to another orbit that had been identified.

To avoid some of these problems in this experiment, Harrison's model (13) for the second zone hole surface of the FS of aluminum was used to calculate the expected calipers for different sample normals and magnetic field angles. The experimental data for a given sample normal yield signals of varying strengths that can be analyzed to obtain plots of caliper versus magnet angle for the given sample. Thus it was desirable for the calculation to yield plots of caliper versus magnet angle plus information about the anticipated
signal strengths at the different angles considered.

To carry out this calculation it is necessary to analyze Harrison's second zone model (Figure 16). The surface is completely determined by fourteen spheres of equal radii and their intersections. So the problem is reduced to a relatively straightforward geometrical exercise. The calculation can be broken into four major areas.

1) Initially the coordinate system \((x,y,z)\) is chosen so the axes are parallel to cubic axes of the crystal. The coordinate system is then rotated using Euler's angles (142, p. 458) such that the new \(z\) axis \((z')\) is parallel to the desired sample normal. This rotation simplified many of the algebraic equations used later in the program to calculate sphere intersections and to allow determination of actual caliper dimensions. In the most complicated part of the program, the rotation reduced many calculations from three dimensions to two dimensions. The new coordinate axes will be labeled \(x',y',z'\).

2) Then fifty-seven (57) equally spaced planes perpendicular to the \(z'\) axis (the sample normal) are passed through the BZ with the central plane always passing through the center of the RZ. The spacing between planes was selected to be large enough that for any possible sample normal the second zone FS would not
Figure 16. The aluminum NPE second zone FS shown with the sample geometry. The coordinate system indicates the coordinates used in the computer calculation following the initial coordinate rotation.
extend beyond either of the outside planes. Each plane is then examined for intersections with the second zone FS.

Clarification of several terms should make the remaining discussion easier to understand. As seen earlier, when a constant magnetic field is applied to a material, the electrons move in $k$-space on constant energy surfaces in planes perpendicular to the magnetic field. The path in $k$-space followed by such an electron around the FS (a constant energy surface) will be called an orbit. In the present geometry with the magnetic field applied perpendicular to the sample normal ($z'$ axis), this means that electron orbits will be in planes parallel to the $z'$ axes and perpendicular to the magnetic field.

Now consider the intersections just found for the second zone FS and the 57 planes of constant $z'$. These intersections are in planes parallel to the magnetic field direction so they do not contain actual paths traversed by the electrons. These intersections simply give the cross section of the FS for a specific value of $z'$ and will be referred to as cross sections. Figure 17 shows three sample $z'$ planes and their resulting cross sections with the second zone FS.
Figure 17. The aluminum NPE second zone FS shown with three typical planes perpendicular to the sample normal as used in step 2 of the computer calculation. The intersections of these planes and the FS result in cross section outlines.
Following the examination of all 57 slices, the information pertaining to each cross section was stored for use later in the program. The number of cross sections found on a single slice varies from zero to six. The information can also be used to plot the cross sections found on each z' slice. For examples see Figure 18, which shows cross section plots for a [100] normal sample for \( z' = 0.00 \), \( z' = \pm 1.3750\bar{A}^{-1} \), and \( z' = \pm 1.4375\bar{A}^{-1} \) respectively.

For \( z' = 0.00 \) the figure is the central cross section of second zone FS. The four curves that the cubic axes pass through are diagonals of the four-sided faces. Each of the remaining four sides represent the common side between two hexagonal faces.

For \( z' = \pm 1.3750\bar{A}^{-1} \) the plane cutting the FS is very near one of the four-sided faces. Note that each of these faces is actually a spherical depression. The plane is chosen to pass between the bottom of the depression and the \( z' \) value where the cross section would break down into four small peaks.

For \( z' = \pm 1.4375\bar{A}^{-1} \) the \( z' \) plane is far enough out the \( z' \) axis that it catches only the four peaks on the second zone FS.

The cross section plots can also be overlaid to give a contour map effect of the second zone hole.
Figure 18. Three typical FS cross section outlines for the NPE aluminum second zone FS shown for a sample with a [100] normal. The distances from the zone center are a) \( z' = 0.0 \), b) \( z' = \pm 1.375 \text{Å}^{-1} \), and c) \( z' = \pm 1.4375 \text{Å}^{-1} \).
surface as seen looking down the z' axis. Sample overlays are shown in Figures 19, 20, and 21 where the sample normals have been chosen as [100], [110], and [111] respectively.

3) Now the cross section information for all the z' slices must be used to obtain caliper information for actual electron orbits. As noted earlier, electron orbits in k-space in a magnetic field are on surfaces of constant energy and in planes perpendicular to the magnetic field. For the problem under consideration, the information on the constant energy surface has already been calculated. Now the direction of the magnetic field must be taken into account.

Since information is desired for calipers as a function of magnetic field angle for ranges of 45° (for a [100] sample normal), 90° (for a [110] sample normal), and 30° (for a [111] sample normal) it is desirable to use an approach that will easily allow the angle of the magnetic field to be changed. This is most easily handled by a simple rotation of the coordinate system around the z' axis so the new y axis (y") is parallel to the magnetic field. The new coordinate system will be x", y", z" where z"=z'=sample normal, y"=direction of the magnetic field, and x" takes its required position. In this new coordinate system,
Figure 19. Cross section overlay plot for the aluminum NPE second zone FS for a [100] sample normal. Cross sections are shown for the plane through the center of the BZ and for every increment of \( z' = 0.0625 \) Å\(^{-1} \) along the sample normal direction. Major symmetry directions are indicated.
Figure 20. Cross section overlay plot for the aluminum NFE second zone FS for a [110] sample normal. Cross sections are shown for the plane through the center of the RZ and for every increment of \( z' = 0.0625 \AA^{-1} \) along the sample normal direction. Major symmetry directions are indicated.
Figure 21. Cross section overlay plot for the aluminum NFE second zone FS for a [111] sample normal. Cross sections are shown for the plane through the center of the BZ and for every increment of z' = 0.0625 Å⁻¹ along the sample normal direction. Major symmetry directions are indicated.
electron orbits in planes perpendicular to the magnetic field are simply in planes of constant $y''$.

In a constant $y''$ plane, points on an electron orbit can be found by evaluating the intersections of the $y''$ plane with the FS cross sections on each $z'$ plane as found in the second part of the calculation. After finding the intersections with all FS cross sections for the 57 $z'$ slices, the resulting set of points describes the actual electron orbits in the $y''$ plane. Figure 22 illustrates a typical $y''$ plane and the intersections found with the cross sections found previously. The resulting orbit is shown in Figure 23 for a [100] sample normal with $\bar{H}$ at 15° from the $x'$ axis and $y''=-0.4375\bar{A}^{-1}$. The caliper for a given orbit is then the largest value of $x''$ between any two points on the orbit. (The use of $x''$ results from the fact that the projection of the real space trajectory on the $y''$ plane is the same shape as the $k$-space orbit but it is rotated by 90° about the direction of the magnetic field, the $y''$ axis. Therefore $x''$ in $k$-space will correspond to the actual $\Delta \bar{r}$ that will span the sample in real space parallel to the $z''$ axis.)

At this point the mechanics have been developed to find the orbit caliper in a given plane perpendicular to the magnetic field. This procedure was then
Figure 22. The NFE second zone FS of aluminum is shown with the lines of intersection with three $z'$ planes. The $y''$ plane is shown perpendicular to the applied magnetic field, $H$. The resulting intersection with the FS indicates an electron orbit in $k$-space.
Figure 23. An electron orbit is shown with its intersections with three \( z' \) planes.

\( z' = z'' = \hat{n} \)

\( y'' = -0.4375 \AA^{-1} \)

\( \theta = 75^\circ \)
repeated for 113 equally-spaced planes of constant \( y'' \) beginning at the center of the Brillouin zone and advancing out the \( +y'' \) axis. The range of \( y'' \) was large enough to be sure that the last planes were beyond the second zone FS for any possible normal and magnetic field directions.

The symmetry of the FS made it possible to determine the required calipers by examining only the \( +y'' \) axis.

4) The remaining problem is to relate these calipers to the intensities expected for RFSE signals. The magnitude of the contribution to the RFSE signal is related to the number of orbits with that caliper.

To calculate this the caliper range from \( \Delta k = 0 \) to \( \Delta k = 4\pi^{-1} \) was broken into 128 bins of equal width. Then the caliper associated with each value of \( y'' \) was examined to find the appropriate bin. The number in that bin was then incremented by one and the procedure was repeated for the next \( y'' \) slice. The end result was that each bin contained an integer representing the number of \( y'' \) slices that had orbit calipers within the range of that bin. This gives information concerning the relative strengths of contributions to RFSE signals for all calipers from \( 0\pi^{-1} \) to \( 4\pi^{-1} \).
This information is valid for only one orientation of the magnetic field relative to the crystal axes. The procedures described in 3 and 4 are repeated for each angle desired between the magnetic field and the crystal axes. The information is then output in tabular form. The vertical axis is the caliper, the horizontal axis is magnetic field angle, and the numbers in the table represent the number of orbits found in that range of the caliper. Figures 24, 25, and 26 show the results for [100], [110], and [111] sample normals.

To aid interpretation, the tabular information can be used to obtain contour plots that show the symmetry and angular trends of the expected data. This is shown in Figures 27, 28, and 29.

The origins of the various lines on the contour plots can be examined by using an option built into the program. It is possible to change one data card and have the caliper for each $\gamma''$ value printed out for each angle examined, so that for each angle a list of 113 calipers is printed. These can then be examined to see from which ranges of $\gamma''$ a signal is resulting. These ranges of $\gamma''$ can then be used with the cross section overlay plots to locate the orbits contributing to the signal. Consider a sample with a [100] normal axis. The signal intensity contour plot in Figure 27 for $\theta=34^\circ$ shows relatively strong signals at Fermi surface calipers of
Figure 24. RFSE signal intensity for NFE second zone FS of aluminum for a [100] sample normal. Numerical values in the table indicate relative intensities. Major crystal axes and their locations are [010] at 0°, [011] at 45°, and [001] at 90°.
Figure 25. RFSE signal intensity for NFE second zone FS of aluminum for a [110] sample normal. Numerical values in the table indicate relative intensities. Major crystal axes and their locations are [110] at 0°, [111] at 35°, [112] at 55°, and [001] at 90°.
Figure 26. RFSE signal intensity for NFE second zone FS of aluminum for a [111] sample normal. Numerical values in the table indicate relative intensities. Major crystal axes and their locations are [101] at 0°, [211] at 30°, and [110] at 60°.
Figure 27. RFSE signal intensity contours for NPE second zone FS of aluminum for a [100] sample normal based on numerical values shown in Figure 24. The initial contour is drawn at an intensity of 2.5 and successive contours at intensity intervals of 3. Major crystal axes and their locations are [010] at 0°, [011] at 45°, and [001] at 90°.
Figure 28. RFSE signal intensity contours for NPE second zone FS of aluminum for a [110] sample normal based on numerical values shown in Figure 25. The initial contour is drawn at an intensity of 2.5 and successive contours at intensity intervals of 3. Major crystal axes and their locations are [110] at 0°, [111] at 35°, [112] at 55°, and [001] at 90°
Figure 29. RFSE signal intensity contours for NFE second zone PS of aluminum for a [111] sample normal based on numerical values shown in Figure 26. The initial contour is drawn at an intensity of 2.5 and successive contours at intensity intervals of 3. Major crystal axes and their locations are [101] at 0°, [211] at 30°, and [110] at 60°.
about $2.70 \times 10^{-1}$ and about $3.00 \times 10^{-1}$. Examination of the computer results indicates that the calipers near $2.70 \times 10^{-1}$ arise from orbits crossing the Fermi surface in the region marked "A" in Figure 30. Similarly the orbits with calipers near $3.00 \times 10^{-1}$ are indicated by the region marked "B" in the same figure. Due to the symmetry of the FS, identical orbits also exist on the other side of the origin from those shown in the figure. This type of orbit location is valuable because it allows examination of the actual orbits contributing to a signal. This will be useful when signals from actual data are compared to these NFE model predictions.

This calculation of calipers is not without limitations. The FS model used does not include rounding effects on the edges and tips of the Fermi surface which result from interactions not included in the nearly free electron model. This will cause all calipers to be too large; however, shapes and relative locations seen on the tabular output and contour plots should still be very helpful.

The calculation does not include any method to handle electron drift velocities that might be helpful in the identification of signals due to non-central orbits. This means that the program cannot be used to examine the consequences of tilting the sample's surface relative to the magnetic field as done for potassium by Peercy et al. (143).
Figure 30. The cross section overlay plot is shown for the MFE second zone FS of aluminum. The magnetic field is indicated. The shaded regions indicate electron orbits that contribute to FS calipers near 2.70Å⁻¹ and 3.00Å⁻¹ that are labeled "A" and "B" respectively.
The calculation examines only full orbits around the FS. It does not yield results relating to the type of partial orbits seen by Gantmakher and Krylov (144) in indium, as discussed earlier.

These limitations should not have too much effect on the usefulness of the caliper results. The trends in calipers as the magnet angle is varied should be very useful. Comparisons can also be made with experimental data using relative magnitudes of calipers at different angles.

The calculation of signal intensities does not take into consideration the shape of the electron trajectory near the sample's surface. Thus the effectiveness of the trajectory as it interacts with the rf field in the skin depth is not considered. Electrons moving nearly parallel to the surface interact with the rf electric field effectively and the longer their path remains in the skin depth, the more they will interact with the field. Trajectories that simply jut into the skin depth and then quickly return to the bulk of the sample are much less effective in their interaction with the electric field. Figure 31 contains examples of effective and ineffective orbits.

For this reason, small changes in intensity as indicated in calculated results may be open to question but large changes are probably a good indication of what to expect in the experimental data. If further information is needed in
Figure 31. Two electron trajectories are shown spanning the sample. The trajectory on the left spends a considerable amount of time exposed to the rf fields in the skin depth and is an effective trajectory. The trajectory on the right spends very little time within the skin depth and is an ineffective trajectory.
this regard, the program can be used (as discussed earlier) to find the actual electron orbits (and trajectories) contributing to the calipers in question. Then the effectiveness of these trajectories can be examined visually to determine if one orbit is expected to be much more effective in contributing to the rf size effect. If so, the calculated intensity results can be expected to show poor correlation to intensities seen experimentally.

This program was written in the PL/1 programming language to allow the use of based variables (145). When using these variables, storage space in the computer is allocated as needed in the calculation. This was very useful in this case because the quantity of information calculated and stored for the cross sections in the beginning of the calculation depended greatly on the sample normal being considered.

Not only quantity but also the organization of this information depended greatly on the normal chosen. This can easily be seen by noting that different $z'$ slices have different numbers of orbits (from zero to six) and orbits have different number of sides (from three to ten). If fixed storage were used, then enough would be required for six orbits with ten sides on each $z'$ slice. This would be extremely wasteful. It would greatly increase running costs and it would also increase turn-around time for the calcula-
By using PL/1, the storage for all this information is exactly the size needed. PL/1 also allows the use of linked lists. This is a convenient method of bookkeeping that allows organization of all the stored items so they can be systematically retrieved for further calculations.
V. THE ALUMINUM EXPERIMENT

A. Apparatus

The RFSE is observed by studying anomalies in the surface impedance of a flat metal sample placed in a static magnetic field. The surface impedance has real and imaginary parts, the surface resistance and surface reactance respectively, and the anomalies can be studied by examining either portion of the impedance. The method of observation used in this investigation utilizes changes in the surface reactance by placing the sample in the system in such a manner that changes in the surface impedance can be readily detected using a standard rf communications receiver. A block diagram of the apparatus is shown in Figure 32.

The sample is placed inside a coil that serves as the inductance portion of a tank circuit used to determine the frequency of a variable frequency oscillator. The coil and sample are then lowered into the liquid helium dewar and located at the center of the pole gap of an electromagnet. Phosphor-bronze springs are used to keep the sample holder assembly centered in the dewar and to decrease vibrations. The impedance of the coil-sample combination reflects both the capacitance and inductance of the coil and the surface impedance of the sample (51). Therefore, when surface impedance anomalies occur, they are reflected in the impedance of the tank circuit and therefore in amplitude and frequency.
Figure 32. Block diagram of the apparatus used in this RPSE investigation. The resulting X-Y recorder trace is $df/dH$ verses $H$. 
changes of the signal in the oscillator.

In all discussions to this point the applied magnetic field has been called a static field. In fact the field only needs to remain constant long enough for an electron to complete its mean free path without its trajectory size changing due to changes in the magnetic field. Since a typical time for an electron between scatterings is about $10^{-14}$ seconds, slow changes in the field will not invalidate any of the principles already mentioned. This result makes it possible to sweep the magnetic field as a function of time and to use a low frequency modulation field which will then facilitate the use of lock-in detection methods for the signal.

Consider the sample and coil subject to a fixed field $H$ plus a small modulation field given by $H_1 \cos \omega_m t$. The frequency of the oscillator can then be written as

$$f(H) = f_\circ(H) + \frac{df}{dH} H_1 \cos \omega_m t + \text{higher order terms in } H_1$$

where $\frac{df}{dH}$ reflects the dependence of the coil-sample impedance on RFSE resonances and magnetoresistance effects in the sample. Cleveland (51) has shown that if the coil has a large quality factor, $Q$, then the value of $\frac{df}{dH}$ is directly proportional to the derivative of the surface reactance with respect to the magnetic field. From this it is seen that the modulation field leads to frequency modulation (fm) of the oscillator frequency with a modulation amplitude of $\left(\frac{df}{dH}\right) H_1$. 
The fm signal from the oscillator is then clipped to generate harmonics. Consider the n-th harmonic of the frequency \( f(H) \),

\[
\text{n-th harmonic} = (n+1)f(H)
\]

\[
= (n+1)f_0(H) + (n+1)(df/dH)H_1 \cos \omega_{mt} + \text{h.o.t.}
\]

where h.o.t. refers to higher order terms. Here it is seen that the modulation amplitude is now given by \( (n+1)(df/dH)H_1 \)
which is enhanced by a factor of \( (n+1) \) over the value seen in the fundamental produced by the oscillator. Thus the signal to noise ratio can be improved by examining the harmonics rather than the fundamental.

The clipper output is detected by an fm communications receiver which is tuned to an integer multiple of the fundamental frequency produced by the oscillator. The output taken from the receiver is chosen to yield information on the modulation amplitude of the input fm signal. This output is directly proportional to \( (df/dH) \).

The receiver output is processed by a tuned amplifier and a lock-in detector (LID) to allow the small signal from the modulation amplitude to be recovered. The LID reference signal is derived from the same oscillator feeding the field modulation coils. The LID output is applied to the Y-axis on an X-Y recorder.

The voltage supplied to the X-axis of the X-Y recorder is taken directly from the magnet power supply which provides
a voltage directly proportional to the value of the magnetic field relative to the end points selected for the field sweep being used. The resulting traces on the X-Y recorder are then plots of \((\text{df/dH})\) versus the magnetic field value.

The individual components represented in the apparatus block diagram will now be discussed and circuit diagrams are given for items designed and constructed during this investigation.

The oscillators and clipper-follower circuits were designed and built for this investigation. The circuits were fabricated on double-sided printed circuit boards with one side serving as a ground plane. Four oscillators were built to cover the frequency range from 3 MHz to 37 MHz. The oscillator circuit design (146) is shown in Figure 33 and the component values are found in Table 1. The inductance \(L\) represents the coil containing the sample located in the liquid helium bath. The oscillators were \#0, \#1, \#2, and \#3 with their frequencies decreasing as their assigned number increased. Oscillators in adjacent frequency ranges overlapped on each end. The exact frequency range of each oscillator for a given sample holder was determined by the area, number of turns, and the wire size used for the sample holder coil.
Figure 33. Circuit diagram for transistor oscillators used in this RPSE investigation. Component values are listed in Table 1. Capacitance values shown on the diagram are microfarads unless indicated otherwise.
Table 1. Components used in oscillators

<table>
<thead>
<tr>
<th>Component</th>
<th>#0</th>
<th>#1</th>
<th>#2</th>
<th>#3</th>
</tr>
</thead>
<tbody>
<tr>
<td>$C_1$</td>
<td>-</td>
<td>-</td>
<td>5</td>
<td>100</td>
</tr>
<tr>
<td>$C_2$</td>
<td>36</td>
<td>36</td>
<td>300</td>
<td>-</td>
</tr>
<tr>
<td>$C_3$</td>
<td>220</td>
<td>27</td>
<td>200</td>
<td>1190</td>
</tr>
<tr>
<td>$C_4$</td>
<td>220</td>
<td>15</td>
<td>200</td>
<td>1290</td>
</tr>
<tr>
<td>$R_1$</td>
<td>4.7K</td>
<td>2.0K</td>
<td>4.7K</td>
<td>4.7K</td>
</tr>
<tr>
<td>$R_2$</td>
<td>10K</td>
<td>12K</td>
<td>10K</td>
<td>10K</td>
</tr>
<tr>
<td>$R_3$</td>
<td>820</td>
<td>680</td>
<td>820</td>
<td>820</td>
</tr>
<tr>
<td>$R_4$</td>
<td>68K</td>
<td>56K</td>
<td>68K</td>
<td>68K</td>
</tr>
<tr>
<td>$R_5$</td>
<td>470</td>
<td>620</td>
<td>470</td>
<td>470</td>
</tr>
<tr>
<td>$L$</td>
<td>2.5</td>
<td>0.5</td>
<td>0.5</td>
<td>2.5</td>
</tr>
<tr>
<td>$Q_1$</td>
<td>2N5245</td>
<td>40822*</td>
<td>2N5245</td>
<td>2N5245</td>
</tr>
<tr>
<td>$Q_2$</td>
<td>2N3904</td>
<td>2N3563</td>
<td>2N3904</td>
<td>2N3904</td>
</tr>
<tr>
<td>$Q_3$</td>
<td>2N3904</td>
<td>2N3563</td>
<td>2N3904</td>
<td>2N3904</td>
</tr>
</tbody>
</table>

* Four lead FET.

All capacitances are in picofarads, resistances are in ohms, and inductances are in millihenries.

The clipper-follower circuit is shown in Figure 34 and was used to generate harmonics of the oscillator frequency to amplify the frequency deviations caused by the RFSE resonance (51). This circuit was mounted on the same panel as the
Figure 34. Circuit diagram for the clipper-follower used in this RFSE investigation. Resistances are in ohms. Capacitances are in microfarads.
oscillators and coaxial cables were used for all interconnections to minimize rf signal losses.

The adjustable Q (quality factor) tuned amplifier was used to increase the signal level and reduce noise at frequencies other than the reference frequency. The design (147) allowed adjustment of both the Q of the filter and the center frequency. Buffer amplifiers were required on both the input and output to avoid a detuning of the circuit by the adjacent circuitry. The circuits used are shown in Figure 35. Initial data were taken using Fairchild uA741 operational amplifiers in the circuit. The signal to noise ratio was later improved when Precision Monolithics operational amplifiers (number Mono OP-05CJ) were used in the circuit. These two different devices are pin compatible so no circuit changes were required to facilitate the conversion.

The receiver used was an Eddystone model 770R (MKII) communications receiver with a frequency range of 19 MHz to 165 MHz. The LID was an Electronics, Missiles, and Communications, Inc. model RJB lock-in amplifier.

A six-inch electromagnet made by Varian Associates was used to provide the "static" magnetic field. The magnet was mounted on a rotating table marked in 10° intervals which allowed rotation of the magnetic field with respect to the fixed crystal. The power supply, a "Fieldial" Mark I, was
Figure 35. a) Block diagram of the adjustable Q, 80Hz tuned amplifier used in this investigation. b) Circuit diagram for the buffer amplifier stages. c) Circuit diagram for the band pass filter stage. Double-ganged 25K potentiometers adjust the center frequency and the 1K potentiometer adjusts the Q. \( C_1 \) is a 20 to 125 pf capacitor. Other capacitances are in microfarads.
used to sweep the magnetic field at a rate that was constant in time (i.e. $dH/dt$ could be chosen and would remain constant during a sweep).

The low frequency signal used to supply the LID reference channel and the modulation field was supplied by a Hewlett-Packard model 204C solid state oscillator. The exact frequency used was chosen to match the frequency of the LID reference filters. The center frequency of the tuned amplifier was then adjusted to the same frequency.

The modulation field was obtained by amplifying the reference signal and applying the resultant voltage to two wire coils wound around the poles of the Varian magnet. The amplifier used to drive the two coils was a single channel Heathkit, model AA-23.

During this investigation two modulation frequencies were examined. All the data to be discussed were taken with 80 Hz modulation. The effect of a lower frequency on the signal to noise ratio was examined by the use of a 42 Hz signal and also a 42 Hz tuned amplifier. No improvement was observed so the system was returned to operation at 80 Hz.

The rf oscillator frequencies were measured using a Monsanto model 110A, programmable counter-timer. When the magnetic field was constant, the oscillators were stable to six significant figures based on a 100 millisecond counting period.
The sensitivity of the apparatus allowed the observation of four NMR signals. The strongest was the NMR due to protons in the GE 7031 varnish used on the oscillator coil. NMR signals were also seen for Cu\textsuperscript{63} and Cu\textsuperscript{65} in the wire of the coil and for Al\textsuperscript{27} in the aluminum sample itself. These NMR signals allowed calibration checks for the magnetic field readings indicated on the "Field dial" magnet power supply.

The design of the dewar system allowed the use of temperatures below 4.2 K. By pumping on the liquid helium the temperature could be lowered from 4.2 K to 1.9 K. No signal enhancement was noted when the system was operated at 1.9 K, but the signal to noise ratio could sometimes be improved this way due to reduced vibration because no bubbles were moving past the sample when the temperature of the helium was below the lambda point.

The detection theory for the fm detection of RFSE signals has been discussed in detail by Cleveland (51).

B. Sample Holder Assembly

A series of sample holder and coil assemblies were made with the idea of making them reusable for different samples. The construction of these assemblies will be discussed and then the technique used to install and remove samples from the coil will be mentioned.

The sample holder assembly was constructed so it could be attached to the bottom end of a coaxial "cable" made of
two sizes of non-magnetic stainless steel tubing, assembled one inside the other. This coaxial "cable" was used to support the sample holder between the pole faces of the magnet to be used. Each sample holder was made from a piece of Synthane rod which was turned down so the two ends of the holder had different diameters. The diameter of the small end allowed a slip fit into the large tube of the coaxial conductor while the large end of the holder was made small enough to fit inside the narrow tail of the helium dewar.

The sample holder was then removed from the lathe and mounted in a jig on an end mill. The jig was designed to guarantee that the axis of the diameters already turned would be perpendicular to the face being cut in this step. If the coaxial conductor holding the sample holder is vertical, then the bottom face on the holder will lie in a horizontal plane.

Holes were drilled in the narrow end and side of the sample holder to allow the electrical connection for the coil to the center conductor in the supporting coaxial conductor.

The coil was preassembled and then attached to the sample holder. The dimensions of the desired coil were chosen based on the dimensions of the sample intended for insertion into the coil. A coil form made of steel shim stock or phosphor bronze was chosen that would provide the proper width and thickness for the coil. The proper length was obtained by varying the number of turns added to the
Once the coil form was prepared, the edges were sanded lightly to remove all burrs that might increase the difficulty encountered when the coil is removed. The coil form was then gripped by one end so the form was perpendicular to the jaws of the vise. This helped ensure that the resulting coil would be rectangular rather than occurring at some skewed angle. The coil form was then coated lightly with vacuum grease. The coil was then wound on the form using the desired gauge of magnet wire with a heavy Polythermaleze insulation. After every two or three turns were added they were pressed snugly toward the jaws of the vise against the preceding turns. This resulted in an even, close-packed coil. After the desired length was reached, the wires were taped to the vise to keep the coil from unraveling and then a smooth-jawed pair of pliers was used to squeeze the large flat sides of the coil against the coil form. This tightened the bends in the wire at the edges of the coil form and resulted in a much flatter coil. One side of the coil was then thoroughly cleaned to remove all vacuum grease and GE 7031 varnish was applied in very light coats. Care was taken to avoid the build up of a thick layer of varnish on this surface. Following a suitable time to allow the varnish to harden, the coil form was pulled loose from the coil to be sure the varnish had not penetrated the coil and vacuum
grease enough to allow the coil to be cemented to the coil form.

The coil was then ready to attach to the sample holder. A table vise was used to press the coil flat against the face of the sample holder until the adhesive dried but special precautions were taken to ensure that the plane of the coil would be perpendicular to the axis of the sample holder. The sample holder was mounted in a machined jig that held the sample holder axis perpendicular to the jaw faces of the vise. Solvents were now used to clean the second side of the coil. Several drops of GE 7031 varnish were placed on the flat face of the sample holder and the clean side of the coil was placed face down in the varnish. The assembly was then placed in the vise and pressure was applied to hold the coil flat against the sample holder face until the varnish dried. It was found that the drying time of the varnish should not be decreased by heating because this caused undesired bubbles in the varnish.

After the varnish dried, the assembly was removed from the vise and all excess vacuum grease was removed. Then new coats of GE 7031 varnish were added to strengthen the bond holding the coil on the sample holder and to add rigidity to the coil itself. The coil form was removed from the coil and the coil form assembly was ready for the insertion of a sample.
It was desirable to be able to insert and remove a sample from a coil without damaging the sample. This led to the use of a thin (0.0004") mylar envelope to transport the sample into and out of the coil. The mylar was cut in a long strip with a width just smaller than the coil width. The strip was then folded with the closed side perpendicular to the long dimension of the strip. The open end of the folded strip was inserted through the coil. Vacuum grease was applied to both inside surfaces of the folded strip near the fold. The sample was then placed inside the folded end. The strip and the sample were then drawn inside the coil. The vacuum grease was used to prevent any looseness in the coil and to hold the sample firmly in the coil. Very severe noise problems in the data were noted when the sample was able to vibrate loosely in the coil.

By using the thin mylar and the sample in the coil, the sample thickness used in a coil had to be less than the thickness of the coil form used. When choosing a coil to use with a sample, the coil form was selected where its thickness was roughly the same as that for two layers of mylar plus the sample. Since the signal strength depends on the fraction of the coil volume that the sample occupies (51), a close fit for the sample-envelope combination in the coil is very desirable.
The sample could be removed from the coil by simply pulling the leading edge of the mylar envelope from the coil. Using this technique, sample Al12 was inserted and removed from coils four times and good signals were still found in each subsequent investigation.

C. Oscillator Problems

In the RFSE the rf oscillator serves two functions. It provides a probe sensitive to surface impedance changes in the sample which can be related to RFSE resonances. Also the frequency of the oscillator determines the skin depth in the metal for the rf fields and hence the width of the RFSE resonance (ΔH) as discussed earlier.

Cleveland (51) has shown that for variable frequency oscillators and the detection method employed in this investigation, the signal represented by the vertical axis on the RFSE data traces is given by

\[
\frac{\text{df}}{\text{dH}} = \left( \frac{n^2 A_s}{4\pi L} \right) \left[ \frac{dX_{xx}}{dH} - \frac{(2r/\omega L)}{dH} \right] \quad (24)
\]

where f=oscillator frequency, \(\omega=2\pi f\), n=turns/length of coil, \(A_s\)=total area of the sample, \(L\)=inductance of the empty coil, \(r\)=effective total resistance of the coil=\text{sum of coil resistance plus effect of sample surface resistance}, and \(X_{xx}\) and \(R_{xx}\) are the imaginary and real parts of the sample's surface impedance. Since \((r/\omega L)^{-1}=Q=\text{quality factor of the coil-sample combination}\), it is seen that for large values of Q the value of \(\text{df}/\text{dH}\) is proportional to \(dX_{xx}/dH\). Equation
24 is valid only for the case where \( L, r, \) and the capacitance of the tank circuit, \( C, \) have no frequency dependence.

Consider the results expected when taking data with two oscillators operating at the same frequency. The parameters \( n, A_0, L, \) and \( r \) are the same for both oscillators since these are all determined by the coil-sample combination and not the oscillator circuit. The remaining parameters, \( \frac{dX_{xx}}{dH} \) and \( \frac{dR_{xx}}{dH}, \) depend only on the sample and the value of the magnetic field so the traces for the two oscillators should have the same shape. But this was not observed in all cases.

Angular data and frequency study data were taken for samples Al 12 and Al 16. Examination of the data traces and the resulting frequency study plots indicated the presence of an unexpected problem. The shape of the background curves differed in some cases and corresponding RPSE signal features were found at different field values for different oscillators. Figure 36 shows traces for oscillators \#0, \#1, and \#2 taken at nearly the same frequency. The presence on each trace of the proton NMR signal due to the GE 7031 varnish on the coil bears testimony to the closeness of the frequencies for each oscillator. Also the NMR signals on each are the same shape although some of the other features seem to be inverted. Notice the differences in the traces from oscillators \#0 and \#1 and the trace from oscillator \#2. Figures 37 and 38 show comparisons of oscillators \#2 and \#3.
Figure 36. Comparison of RFSE data traces taken with oscillators #0, #1, and #2. Experimental conditions were identical for all traces.
Figure 37. Sample traces for oscillators #2 and #3 for identical experimental conditions.
Osc #2
f = 19.174 MHz

Osc #1
f = 19.199 MHz

SAMPLE NORMAL = [111]

H AT 15° FROM [110] AXIS

Figure 38. Sample traces for oscillators #1 and #2 for identical experimental conditions
and oscillators #1 and #2 for a smaller range of magnetic field. Again note the differences in the traces from oscillators #1 and #2. The traces from oscillators #2 and #3 show great similarity.

The problem is also evident if data for a frequency study are recorded over a frequency range requiring the use of all four oscillators. The result of such a study made on sample Al 16 as shown in Figure 39 where points resulting from oscillators #0 and #1 are represented by the symbol "x" and points from oscillators #2 and #3 are represented by the symbol "+". It is seen that the sets of points, "+", form reasonably straight lines but they do not line up with the points from oscillators #0 and #1. This is extremely evident in the region of frequency overlap for the two sets of data.

One explanation considered was the possibility that the high frequency data points were actually at higher values of frequency than the Monsanto counter indicated, perhaps two or three times the measured values. If this were true, the shift to lower field values would result from the natural narrowing of the skin depth with increasing frequency, the very effect that leads to the frequency study concept. The waveforms of the signals leaving the oscillators were examined with a wide band oscilloscope and found to be compatible with the frequency counter readings. The signals were not pure sine waves but it was not possible to see if the distor-
Figure 39. Frequency study plot for Al 16 with data from oscillators #0 and #1 ("x") and #2 and #3 ("+")
tion was actually present on the coil or if it was introduced in the follower circuit mounted on the oscillator circuit board. The magnetic field values for the proton NMR signals also agreed with the frequency readings.

Variations in the magnitude of the rf fields at the sample were considered. Additional resistance was added to the coaxial line between the oscillator and the coil to decrease the rf currents flowing in the coil. Data traces were taken and compared as each increment of resistance was added. No effect was seen on RFSE signal locations or the background signals. The signal to noise ratio decreased as resistance was added until the oscillations finally ceased.

Adding resistance to the line would increase the value of \( r \) in Equation 24 and affect the value of \( \frac{df}{dH} \) if the second term in that equation was noticeable for these experimental conditions. No such effects were noted.

The clipper circuit was examined to determine if it could be leading to the problem in some unexpected manner. Signals were recorded using four other clipper designs with no change observed in signal locations. Again the only effect was on the signal to noise ratio. The signal was also recorded without a clipper and the same signals were found again. This was true even when the receiver was tuned to the tenth harmonic of the oscillator's fundamental frequency. This indicated either a non-linearity in the signal process-
ing by the follower on the oscillator board itself or the presence of harmonics in the actual oscillator circuit. The search for harmonics in the oscillator circuit was directed at observing the proton NMR resonance for harmonics of the oscillator's fundamental frequency if these harmonics did in fact reach the coil.

The first evidence of actual harmonic content at the coil was observed for tube oscillator A where a proton NMR signal was observed at a magnetic field corresponding to three times the fundamental frequency of the oscillator. This oscillator had been used in an earlier RFSE investigation (51, Cleveland oscillator #3) and gave RFSE signal traces in this investigation that were the same as those for oscillator #1. In Figure 40 the upper trace was taken with oscillator A for the magnetic field at an angle which yielded good RFSE signals. The lower trace was taken for a magnetic field direction that maximized the NMR signals and minimized the RFSE signals. The proton NMR signal is seen in response to the fundamental \( f_o \) and its second harmonic \( 3f_o \). From Figure 40 it is seen that the \( \frac{df}{dH} \) signal due to the fundamental NMR signal is roughly seven times as strong as that due to the second harmonic.

Similar examinations were carried out to evaluate the harmonic content in the signals obtained using transistor oscillators #1, #2, and #3. It was impossible to examine
Figure 40. Sample data traces illustrating the presence of the second harmonic found in the signal from tube oscillator A.

SAMPLE NORMAL = [111]
TUBE OSCILLATOR A
f₀ = 12.819 MHz

df/dH

PROTON NMR (f₀)

PROTON NMR (3f₀)

H 15° FROM COIL AXIS

11 COIL AXIS
AMPLITUDE + BY 2.5

MAGNETIC FIELD (kOe)

1.0 3.0 5.0 7.0 9.0 11.0
oscillator #0 due to the limited range of magnetic field supplied by the electromagnet. The results of this survey will be summarized below.

Oscillator #3 showed no indication of the second harmonic NMR signal. NMR signals were seen due to Cu$^{63}$, Cu$^{65}$, and Al$^{27}$. The amplitudes of these signals as measured directly from the data sheets were about 7%, 3%, and 8% of the strength of the fundamental proton NMR signal.

Data traces obtained using oscillator #2 indicated a proton NMR signal due to the second harmonic of the measured frequency. This signal had an amplitude of about 2% of the fundamental proton signal. The NMR signals due to Cu$^{63}$ and Al$^{27}$ were also seen with this oscillator and were a little stronger than the second harmonic peak.

Oscillator #1 was examined and no harmonic content was noted. This investigation was limited by the field range of the electromagnet used.

A brief study was made with tube oscillator A operating between 13.1 MHz and 15.3 MHz and large changes in the harmonic content were noted. These changes were roughly an order of magnitude as measured on the data traces. Caution should be used when considering these values. The apparatus used in the fm detection method is not designed to retain exact amplitude information for the signals, and amplitude fluctuations are often seen in traces taken under the same
experimental conditions. Even so, the wide range of amplitude seen here can be considered to reflect a real trend in the signal's harmonic content.

Another attempt to evaluate the harmonic content of the signal on the sample holder coil utilized an external pick-up coil and an auxiliary rf signal generator. The pick-up coil was placed in close proximity to the sample holder coil (just outside the glass dewar) and the coil output was fed to the communications receiver input. The signal strength as indicated by the receiver S-meter was recorded with the receiver tuned to each harmonic value. The rf generator was then tuned to the same frequencies and the signal level that gave the same signal strength was recorded. This study indicated that oscillator #3 applied the purest sine wave to the sample holder coil. Tube oscillator A and transistor oscillator #0 showed the greatest amount of harmonic content on the sample holder coil. Oscillator #2 exhibited less harmonic content than #1. Problems arose in these measurements associated with resonances within the probe coil, local radio station pick up, and impedance matching of the coil to the receiver.

Following these studies oscillators #0 and #1 were discarded and oscillators #2 and #3 were used to take angular data on the aluminum samples. The reasons for this decision are given below.

1) The large harmonic components found in oscillators #0
and A and the similarity of their data traces to those obtained using oscillator #1.

2) The inconsistencies in the high frequency data points on Figure 39. The points taken with #0 and #1 do not show the standard straight line form expected.

3) The very small harmonic content found in oscillator #3 and the similarity between its traces and those for #2.

4) The data for #2 and #3 show good alignment on the frequency study shown Figure 39.

From the data recorded, it is seen that the f₀ and 3f₀ signals do not behave as two independent signals affecting the coil-sample system in a linear manner. Clipping a signal of frequency f₀ generates harmonics at 2f₀, 3f₀, 4f₀, etc. Clipping for a signal at 3f₀ results in harmonics at 6f₀, 9f₀, 12f₀, etc. For a linear superposition of these signals, both signals will be present for very few frequencies that could be tuned on the communications receiver (e.g., 6f₀, 9f₀, etc.). But experimentally it was found that both NMR signals could be observed when the receiver was tuned to adjacent harmonics of f₀. This indicates that for the interactions taking place in this experiment, the idea of a linear superposition of responses to the signal on the coil is not valid.
D. Sample Preparation

All samples prepared were taken from a rectangular parallelepiped (16 mm x 17 mm x 65 mm) of 99.9999% pure aluminum obtained from Cominco American, Inc. of Spokane, Washington. The residual resistance ratio (the resistance at 300 K divided by the resistance at 4.2 K) quoted by the manufacturer was 13,000. (After spark cutting had been completed a thin needle of aluminum was examined using a standard four probe technique and the resistance ratio was found to be greater than 8500. Greater accuracy could not be obtained due to small magnitude of the signals being measured and the noise level in the measuring system.) The ingot was not a single crystal but after chemical etching in a mixture of phosphoric, sulfuric, and nitric acids (148), large crystallites were visible to the naked eye. The last 20 mm of material on one end of the ingot appeared to be a single crystal. That end of the ingot was then examined using Laue x-ray backscattering techniques. This was done to locate the three cubic axes in the crystal so cuttings could be planned to minimize crystal wastes when [100], [110], and [111] normal samples were to be prepared.

For each orientation of samples to be cut the crystal was aligned to within 1° of the desired axis using the x-ray techniques. Several thin plates (1 mm thickness) were then cut off the end of the ingot with a moving-wire spark cutter.
Extra plates were cut in each case to allow the preparation of five or six RFSE samples for that orientation before further cutting from the ingot would be required. Each slice was then re-etched in the acid solution to allow a visual check for any crystal boundaries in the face of the slice. Each slice was then spark cut into rectangular plates with surface dimensions of roughly 8 mm x 9 mm. These rectangular plates were then prepared as individual samples. For further preparation of the sample's first side, the sample was attached to an adjustable lapping plug as discussed by Cleveland (51) and x-ray techniques were used to align the crystal to within less than 0.5° of the desired axis.

For use in the RFSE, the samples need to have very flat, parallel faces and be as free of damage as possible. Therefore a method of sample preparation would be preferred if it would not induce large strains in the crystal. For this reason, the technique of Spong and Kip (31) was tried. This involved a modified electropolishing process which incorporated a lapping motion against a Teflon cloth submerged in the acid solution. The sample was mounted to an aluminum lapping jig using a standard mixture of graphite and Duco cement. The portion of the jig surrounding the sample was then coated with a lacquer to limit current flow to the sample surface. The lapping jig acted as the anode for the electropolishing and the cathode was a sheet of stainless
steel mounted below the Teflon lapping cloth. Very flat surfaces resulted from this technique, but the technique was discarded when it was found that the sample's second surface could not be prepared without severe damage occurring on the first side that had already been prepared. This resulted because the acid solution could not be prevented from seeping behind the crystal and reetching the first side. This problem was compounded by the smooth surface on the first side and the fact that the Duco cement and graphite did not adhere well to the finished surface. Different mixtures of the conducting adhesive were tried as well as a silver paint but none were successful for holding the sample down and preventing the acid from etching the first side.

Good samples were then prepared using mechanical lapping techniques. Samples were attached to the adjustable lapping plug using beeswax. Lapping was done in four steps. The first employed #600 grade grit paper and methanol as a slurry. This was used until all signs of roughness due to spark cutting were removed and the sample thickness was reduced by about one third. Next a 4/0 emery paper was used to remove all signs of scratches left by the #600 grade grit. Then one micron alumina abrasive was used with a water slurry for lapping on a Teflon cloth. This was continued until the surface had a uniform texture when viewed under a low power microscope. The surface at this point was not
shiny but rather had a smoky appearance. The final lapping was done using a commercial red rouge with methanol on a Teflon cloth. The size of the abrasive particles in the rouge were less than 1/3 of a micron in diameter. Following this the surface appeared very flat and shiny to the naked eye. Some small scratches could be seen under the microscope but the surface was generally very nice. Care was taken to keep the alumina abrasive from being transferred to the Teflon used for the final lapping.

The sample was removed from the lapping plug and the plug face itself was then lapped flat to ensure that it was perpendicular to the axis of the plug. The sample was then reattached to the plug so the unfinished surface could be prepared. Before any work was done on the surface the crystal was x-rayed again to check alignment. This was done to guard against the chance of any foreign particles in the adhesive getting under the crystal and preventing it from being flatly held against the plug face.

The second face was then prepared following the same steps used for the first side. The only difference being that the #600 grade grit paper was used until the desired sample thickness was approached. Thickness was monitored during preparation using a Leitz optometer.

The sample was then thoroughly cleaned with warm solvents (petroleum ether, acetone, and trichloroethane) to
ensure a uniformly clean surface for dip electropolishing. A standard electropolishing solution of 6% perchloric acid in methanol was used in a dry ice and acetone bath. The current density was set in compliance with the guide lines set forth by Metz (149). The experimental findings of Metz make it possible to monitor current and voltage changes to avoid current densities that will probably lead to pitting rather than a smooth shiny surface.

Following the electropolishing, the surfaces were shiny and flat but showed signs of rounding near the edges. In some samples a few scratches were visible under the microscope but the majority of the surface was clear of such obvious defects. Further x-ray photographs taken at this time showed sharp distinct spots with no signs of twinning or obvious crystal damage resulting from the surface preparation.

E. Sample Thickness

The thickness of the samples has been determined by two methods. The first involves measuring the mass of the sample using an accurate analytical balance. The calibrated dimensions of the sample are then measured using a calibrated traveling microscope. The linear dimensions of the surface are then used to calculate the area of the sample. The average thickness was then calculated by using the known density for aluminum (2.702 gm/cm³). These values are shown in the third column of Table 2. This procedure includes any
rounding near the edges into the average. Thus this technique may yield an average that is lower than the average away from the edges if rounding problems are important for a specific sample.

Table 2. Average thickness for aluminum samples

<table>
<thead>
<tr>
<th>Sample</th>
<th>Normal axis</th>
<th>Average thickness (by mass) (microns)</th>
<th>Average thickness (by microscope) (microns)</th>
<th>Microscope readings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al 12</td>
<td>[100]</td>
<td>172.3</td>
<td>172.3± 4.9</td>
<td>36</td>
</tr>
<tr>
<td>Al 13</td>
<td>[100]</td>
<td>150.0</td>
<td>149.7±10.6</td>
<td>39</td>
</tr>
<tr>
<td>Al 7</td>
<td>[110]</td>
<td>180.9</td>
<td>188.4± 7.8</td>
<td>36</td>
</tr>
<tr>
<td>Al 16</td>
<td>[111]</td>
<td>112.5</td>
<td>110.8± 7.8</td>
<td>48</td>
</tr>
</tbody>
</table>

The second method involves taking a series of actual thickness measurements across the face of the sample using the focal point of a microscope with a calibrated lens movement. The sample is first placed on a flat metal plate and held in place with beeswax. The sample was then measured using a microscope with a calibrated table motion in two directions perpendicular to the axis of the lens system. This made it convenient to take data at equally spaced points across the surface. Thickness measurements were recorded at
the points on a square grid where the grid lines were separated by 0.050 inches in both directions. The recorded values were averaged and the standard deviation was calculated for each sample. Figure 41 contains detailed information about the distribution of the thickness measurements for the four samples of importance in this experiment.

The average thicknesses and standard deviations for the samples are listed in column four of Table 2. The number of readings taken for each sample is recorded in the last column of the same table.

It should be noted that excellent agreement in the average thickness resulting from the two techniques is found for samples A1 12, A1 13, and A1 16. The result for A1 7 shows agreement within the indicated standard deviation. The disagreement is also compatible with effect of severe rounding near the edges of the sample.

None of the samples are as flat as desired. On the type of graph seen in Figure 41, a flat sample would find all reading of the thickness at the same point. A wedge-shaped sample would lead to a graph with an even distribution of values from the minimum thickness to the maximum thickness.
Figure 41. Distribution of thickness measurements for samples Al 12, Al 13, Al 7, and Al 16.
VI. DATA AND DISCUSSION

During the course of this experiment angular and frequency study data were recorded for a series of samples with [100], [110], and [111] axes normal to their respective sample surfaces. The following discussion will deal with the raw angular data, the frequency study data, and the resulting angular dependence of the calipers and the electron orbits contributing to the individual signals in that order.

A. Raw Angular Data

The angular data considered were taken using samples Al 12 ([100] normal axis), Al 7 ([110] normal axis), and Al 16 ([111] normal axis). The symmetry of the FS then requires that the RFSE signals show 90°, 60°, and 180° rotational symmetry for the respective samples. This can easily be seen by examining the FS cross section overlay figures for the normal directions (Figures 19, 20, and 21).

The Al 12 ([100] axis) data and Al 16 ([111] axis) data did not demonstrate the proper angular symmetry. First consider the data obtained using Al 12.

1. [100] sample normal

The initial data were recorded with one of the cubic crystal axes parallel to the coil axis. A rectangular plot of the uncorrected field values versus the magnetic field angle is shown in Figure 42. Data are shown for every feature on the data traces and not simply the lowest feature of
Figure 42. Angular dependence of raw angular data for [100] sample normal with the coil axis along a cubic crystal axis.
every signal. Data are not shown for negative field angles because they were simply a mirror image of the data shown.

From the contour overlay plot (Figure 19) it is seen that the data should possess both 90° rotational symmetry and mirror symmetry about the [010] and [011] axes in the plane of the sample. The mirror image about the [010] axis coinciding with the coil axis did occur but there was no mirror imaging about the [011] axis or any 90° rotational symmetry.

Some apparent deviation from the required symmetry could be expected due to the decrease of experimental sensitivity as the magnetic field approached 90° on the figure, but this can not explain the deviations from mirror symmetry to be seen in the region of the [011] axis. In this small angular range the sensitivity should not change greatly. Examining the features in this region for field values near 600-700 Oe and near 1000 Oe shows that they reach their extrema about 51°-52° from the [010] axis and not at 45° as expected.

The sample was removed from the coil and then replaced in the coil after a 90° rotation. This placed the other cubic axis in the plane of the sample parallel to the coil axis. Angular data were recorded again and the features were unchanged from those seen in Figure 42.

The sample was again removed from the coil. A new coil was prepared that allowed the sample to be placed inside the
coil with a [011] axis parallel to the axis of the coil. Due to space limitations in the dewar system, this coil did not completely enclose the sample as the previous coils had done. It simply belted the sample, passing across two opposing corners of the rectangular sample. The coil width was adequate to cover more than one-half of the sample's area.

The resulting data are shown in Figure 43. Two striking features are to be noted. First, many of the features seen earlier near 0° are now seen at 45° as expected. This is most obvious for features near 1000 Oe. Second, some features still do not demonstrate the expected symmetry and these features are again symmetric about the coil axis.

2. [111] Sample normal

The symmetry problems for sample Al 16 (a [111] normal axis) are not as pronounced as those just examined. Much of the apparent problem here may result from the complexity of the signals seen near the [101] axes. Figure 44 shows the data for Al 16 with the [101] axis 7° ±1° from the coil axis. To resolve the data near the [101] axis, data were taken at much closer intervals than for the remainder of the angular range. Fourteen angles were examined in the range from 0° to ±10° and then 2° intervals were used outside that region. The extreme care taken near the [101] axis was required because the features on the traces were changing very rapidly as the angle was varied. To avoid misinterpretation of the
Figure 43. Angular dependence of raw angular data for [100] sample normal with the coil axis at 45° from a cubic crystal axis.
Figure 44. Angular dependence of raw angular data for [111] sample normal with the coil axis 7° from the [101] axis
data, the angular intervals were decreased to allow accurate
determination of the behavior for individual features.

Some features near 60° (the adjacent [\( \bar{1}10 \) axis)
continue rising to higher field values beyond the ±60° lines.
This disagrees with the symmetry required. Interpretation
of this region is complicated by the weakened signals due to
the angle between the coil axis and the applied magnetic
field and the complexity of the equivalent region seen near
0°.

Another symmetry problem exists near the [11\( \bar{2} \)] axis
where mirror symmetry is required for the data. The small
peak near 30° and 1100 Oe does not demonstrate mirror symme-
try. This feature occurs at larger field values for 0 above
30° than it does below 30° for equal distances from the 30°
line. This again is evidence of a symmetry problem in the
data.

The sample was removed from the coil and reinserted
after a 90° rotation in an effort to obtain data
complementing that seen in Figure 44. This is the same pro-
cedure that was followed for the [100] sample as previously
discussed. A plot of the data obtained is shown in Figure
45. Notice that the general features of these data do show
symmetry under a 60° rotation as required, but the feature
near 0° ([11\( \bar{2} \)] axis again) and 1100 Oe now does not show
mirror symmetry about the 0° line.
Figure 45. Angular dependence of raw angular data for [111] sample normal with the coil axis 30° from the [101] axis.
3. [110] sample normal

The raw data from Al 7 (a [110] sample normal) followed the symmetry pattern expected. The data therefore presented no unusual problems and will not be discussed here. The resulting calipers will be discussed later in the text.

4. raw data discussion

From the raw data presented above, the worst symmetry problems occurred for Figure 42 where the data for a [100] sample with a cubic axis along the coil axis were presented. This figure showed unexpected symmetry for all features above 500 Oe. The data for the same sample following a 45° rotation inside the coil (a [011] axis along the coil axis), (Figure 43) showed the proper symmetry for all features above 1000 Oe for angles that gave reasonable sensitivity for the RFSE signals (ie. ±75° from the coil axis). Below 1000 Oe some of the data does not follow the expected pattern at angles greater than ±45° from the coil axis.

The problems with the [111] data were considerably less. The complexity of the signals near the [101] axes could easily result in misinterpretation of data traces for magnet angles where the RFSE sensitivity is diminished. Thus the only real question relates to the lack of mirror symmetry about the [112] axis for the feature near 1000 Oe.

It should be noted here that for the data presented in Figure 44, the coil axis and the [101] axis in the sample
were $7^\circ \pm 1^\circ$ from each other. Thus if data depending only on the angle between the coil and the crystal axis occurred here, their symmetry axis should be different than the symmetry axis for the RPSE data resulting from the FS. This is not seen in the Al 16 data.

Asymmetry in the data could have been caused if the magnetic field was not applied parallel to the sample's surface. This could give rise to data with $180^\circ$ symmetry if the signals were affected by the tilting. Several measurements were made to examine this possibility.

The magnet assembly was examined to determine if its axis of rotation was along a vertical axis. If the magnet base plate had been installed poorly this problem could have arisen. The deviation from the vertical axis was found to be $0.60^\circ$. The deviation was in a direction that would cause the effects of the tipping to be symmetric about magnetic field angles of $120^\circ R$ or $150^\circ L$ as seen on the magnet base plate. The problems of data symmetry are not compatible with these findings as their symmetry axes occurred far from either of these axes.

Data were also taken with the top of the sample holder displaced horizontally from its normal position. This motion caused the stainless tubing supporting the sample to be tilted slightly. The sample would tilt a similar amount. This led to the sample being tilted $\pm 0.7^\circ$ about two perpen-
dicular axes. No effect of any kind was observed in the data traces recorded.

Based on these two studies it is felt that the symmetry problem did not arise from an inclination of the magnetic field relative to the plane of the samples.

Data exhibiting a similar lack of proper angular symmetry has recently been reported by Matthey et al. (124) in white tin. In their data and the data presented here the signals not agreeing with the crystalline symmetry are symmetric about the coil axes. One possible explanation is the unknown importance of the polarization of the rf fields incident on the sample's surface relative to the crystal axes in the sample. The solution may also be the angular relationship between the modulation field and the coil axis. This has been considered and no mechanism has been visualized that would allow the modulation field information to feed through the system except the RFSE.

The symmetry problem in the data of Matthey et al. (124) occurred for a sample with a [001] sample normal (the axis in white tin for four fold symmetry). Examining their data and the data presented here have led to one condition in the two cases that is similar. The tin data with unexpected symmetry were recorded when the magnetic field was rotated in a plane perpendicular to the axes of third zone pieces of the FS shaped like cylinders. Therefore, the rf electric field was
always perpendicular to these cylinders. In their other data, the sensitivity to certain signals depended greatly on orientation of the rf field relative to these cylinders.

In the case of aluminum, the third zone FS is similar to four cylinders connected to each other around each square face of the BZ. The worst symmetry problem seen in the data presented here (Figure 42) occurred when the rf electric field was perpendicular to two of these square face assemblies. In the remainder of the data taken the rf electric field was never perpendicular to any entire assemblies. In some cases it was perpendicular to two of the four segments on two faces (Figures 43, 44, and 45 and the [110] data). Presently no mechanism is visualized to explain this problem based on the orientation of the rf fields relative to these cylinders. It is apparent from studying the data that two processes are interacting to yield the observed data. The process leading to the symmetry problems is currently unexplained, but by taking complementary data the RFSE signals can be extracted.

In the raw data presented, the existence of complementing data resulting from sample rotations inside the coils made it possible to examine two sets of data for consistent data for both the [100] and [111] samples. This comparison of data made it possible to reject signal features that seem to result from the relationship between the coil
axis and the crystalline axes. It was also possible to eliminate the line "A" at 500 Oe in Figure 42 because this feature did not scale with changing sample thickness when preliminary data for another [100] sample were examined.

The data remaining after this examination have been used to determine FS calipers and will be discussed in a later section.

B. Frequency Studies

Frequency study data were recorded for several samples with [100] and [111] sample normals. The only data showing the frequency dependence of Equation 16 were obtained using Al 13 (a [100] sample normal). The data recorded for Al 12 showed erratic frequency dependence as will be seen later. These facts will be examined using the ideas discussed in an earlier section relating to frequency study problems.

Consider the data for two samples with [100] sample normals, Al 12 and Al 13. Recall from the discussion of samples with non-uniform thickness that the low field edge of the RFSE signal arises from the thickest part of the sample. From Figure 41 it is then seen that the onset will occur for t=180 microns for Al 12 and for t=164 microns for Al 13 (ignoring the isolated point at 167 microns). Thus the corresponding features for Al 13 should be at field values about 10% higher than their counterparts for Al 12. This is seen to be true for the two traces shown in Figure 46 which were
Figure 46. Data traces for Al 12 and Al 13 for the same crystalline direction using oscillator #2.
taken for the same orientation of the magnetic field relative to the crystal axes.

Next consider the thickness distribution for the two samples as seen in Figure 41. It is seen that Al 12 has a smaller variation in thickness but also a more erratic thickness distribution than seen for Al 13. The average and the standard deviation of the number of microscope readings per micron for every value of thickness between the minimum and maximum values have been calculated for these samples. The values for Al 12 and Al 13 are $1.89 \pm 1.56$ and $1.06 \pm 0.98$ respectively. This means that Al 13 is more wedge-shaped than Al 12 and the behavior of the RFSE signals should be examined with this in mind. Consider feature "A" on the Al 13 trace in Figure 46 and the simplified signal behavior seen for a wedge-shaped sample in Figure 11b. Notice in Figure 11b the first and last features are the same shape as each other but are inverted with respect to each other. This is the general behavior exhibited by feature "A" for Al 13. This trace also shows great structural similarity to the Gantmakher (60, Figure 19) illustration of RFSE line splitting resulting from a wedge-shaped sample. Next consider feature "A" for Al 12 and the curve seen in Figure 12b. Feature "A" for Al 12 is seen to show more structure in its central region than its counterpart from Al 13. This is exactly the same behavior as seen when comparing Figure 12b for a
wavy sample with Figure 11b for a wedge-shaped sample.

If Al 13 is behaving like a wedge-shaped sample then the upper and lower features of signal "A" should result from the thinnest and thickest portions of the sample and the field values should be scaled accordingly. This can be expressed as

$$\frac{B_{\text{max } t}}{B_{\text{min } t}} = \frac{t_{\text{min}}}{t_{\text{max}}}.$$  \hspace{1cm} (25)

The left hand side of Equation 25 can be evaluated from the RFSE data and the right hand side can be evaluated with the aid of Figure 41. From Figure 46 it is seen that

$$\frac{B_{\text{max } t}}{B_{\text{min } t}} = \frac{1138}{1296} = 0.88$$ \hspace{1cm} (26)

and from Figure 41 it is seen that

$$\frac{t_{\text{min}}}{t_{\text{max}}} = \frac{136}{164} = 0.83.$$ \hspace{1cm} (27)

The field values were chosen as two corresponding features on the line shape of signal "A". The onset of the signals was not used in an effort to minimize any problem caused by the interaction of the two signals in the central region of the resonance. This numerical agreement is satisfactory in light of the unknown interaction of the two signals within feature "A".

Now consider the frequency study data taken for these two samples. Frequency study plots are shown in Figure 47. The data for Al 12 show no systematic frequency dependence. The lines for Al 12 have been examined using a linear least-squares fit to the mathematical form of Equation 16. The
Figure 47. Frequency study plots for Al 12 and Al 13 for the same crystalline direction using oscillator #2
value of the exponents, \( \alpha \), giving the smallest chi-squared for the five lines were 0.09, 0.07, 1.61, 0.17, and greater than 2.4 starting with the lowest line of Figure 47a. This behavior is not consistent from line to line and the data have been considered to be of no value for a useful frequency study. The data for Al 13 examines the frequency dependence of feature "A" on Figure 46. Examine lines (1 and 2) and (4 and 5). These lines correspond to features presumed associated with the RFSE signal arising from the thickness extremes. Line 3 will be ignored to avoid possible problems in the central region of the signal where the two signals might be interacting. The data for lines 1, 2, 4, and 5 were examined for compatibility with the frequency dependence expressed in Equation (16) using a linear least-squares fit program and the Cleveland method for determining proper field values. Both of these techniques have been discussed in detail in a preceding section.

The linear least-squares examination of the data led to the information found in Table 3. The first two columns give the line number and the value of the exponent, \( \alpha \), corresponding to the smallest value of chi-squared for each line. The exponents for the two lines in each pair are in good agreement. For each pair of lines a value of the exponent was selected to represent the real exponent for the two lines, \( \alpha(\text{real}) \). These values and the corresponding line
intercepts and slopes are given in the last three columns of the table. These data for Al 13 show good frequency dependence for the signals arising from both sample thickness extrema. This is to be expected since at each extremum for a wedge sample, only the skin depth can change to affect the width of the RFSE signal.

Table 3. Al 13: Linear Least squares fit results

<table>
<thead>
<tr>
<th>Line</th>
<th>Best fit exponent</th>
<th>Selected real exponent</th>
<th>Intercept (Oe)</th>
<th>Slope Oe (MHz)(^{1/3})</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.46</td>
<td>0.48</td>
<td>1125</td>
<td>73</td>
</tr>
<tr>
<td>2</td>
<td>0.51</td>
<td>0.48</td>
<td>1160</td>
<td>128</td>
</tr>
<tr>
<td>4</td>
<td>0.49</td>
<td>0.46</td>
<td>1269</td>
<td>95</td>
</tr>
<tr>
<td>5</td>
<td>0.45</td>
<td>0.46</td>
<td>1310</td>
<td>147</td>
</tr>
</tbody>
</table>

The results of the Cleveland technique for finding the proper exponent for each pair of lines are 0.19 for lines 1 and 2 and 0.17 for lines 4 and 5. The corresponding intercepts for two sets of lines are 1082 Oe and 1201 Oe respectively.

The numerical results for these frequency study data can now be analyzed using the ideas previously developed for wedge-shaped samples. The comparisons should be realistic.
because the frequency range for the data generated for the earlier discussion is identical to the frequency range for the Al 13 data. First consider the ratio of the "Cleveland best fit exponent", $\alpha_{\text{min}}$, to the selected real value of the exponent for each set of lines, $\alpha_{\text{real}}$. For lines 1 and 2 this yields 0.40 and for lines 4 and 5 it yields 0.37. These values can be used with Figure 13 to obtain an indication of the relative importance of skin depth broadening and thickness variation broadening for the sample. From Figure 13 it is seen that the value of RATIO deduced for lines 1 and 2 is between 0.62 and 0.81. Similarly for lines 4 and 5 the range for RATIO is 0.70 to 0.91.

RATIO can also be found directly from the lines of the frequency study as discussed earlier. Recall that RATIO is defined as the largest spread in intercepts for a family of lines divided by the largest difference in slopes for the same lines. These values are taken from the lines when they are plotted on an axis system based on the real value of the exponent for the family of lines. Using this definition it is seen that for lines 1 and 2,

$$\text{RATIO (1 and 2)} = \frac{35}{55} = 0.64.$$  \hspace{1cm} (28)

Similarly for lines 4 and 5,

$$\text{RATIO (4 and 5)} = \frac{41}{52} = 0.79.$$  \hspace{1cm} (29)

Both of these values are seen to fall in the anticipated range found from Figure 13. It is also worth noting that
based on the values of \( \alpha_{(\text{min})}/\alpha_{(\text{real})} \) for these two sets of lines and Figure 13, the value of RATIO (1 and 2) is expected to be less than RATIO (4 and 5). This is seen experimentally.

Another comparison between the experiment and the theoretical predictions on this matter can be made concerning the amount that the Cleveland method reduces the intercept for a family of lines. Using the data on the individual lines from Table 3 and the common intercepts for the two series of lines for their respective values of \( \alpha_{(\text{min})} \) it follows that

\[
(A(1, \alpha_{(\text{real})}) - A(\text{ave}) (1\text{ and }2, \alpha_{(\text{min})})) = 1125-1082=43 \text{ Oe} \quad (30)
\]

and

\[
(A(4, \alpha_{(\text{real})}) - A(\text{ave}) (4\text{ and }5, \alpha_{(\text{min})})) = 1269-1201=68 \text{ Oe}. \quad (31)
\]

The theoretical predictions can be based on the spread in the intercepts for the two series of lines for their appropriate values of \( \alpha_{(\text{real})} \).

Intercept spread (1 and 2) = 1160-1125=35 \text{ Oe} \quad (32)

Intercept spread (4 and 5) = 1310-1269=41 \text{ Oe}. \quad (33)

The value of \((A(\text{real})-A(\text{min}))\) is expected to be roughly equal to the spread in intercepts for the lines as discussed earlier. Thus it would be expected that for lines 1 and 2 the intercept will be reduced about 35 Oe and for lines 4 and 5 the reduction will be about 41 Oe. From these two cases it is seen that the qualitative agreement is good. The ratios of the experimental to theoretical reductions in the intercepts for the two sets of lines are:
Lines 1 and 2: \((\frac{43}{35}) = 1.23\) \hspace{1cm} (34)

Lines 4 and 5: \((\frac{68}{41}) = 1.66\). \hspace{1cm} (35)

In both cases the reductions were larger than expected. The largest reduction was predicted and observed for lines 4 and 5 rather than for lines 1 and 2.

The theoretical concepts examined earlier for wedge-shaped samples have been examined as they relate to experimental frequency study data. The idea of signal splitting due to a wedge-shaped sample has been substantiated experimentally. It has also been noted that each of these split signals demonstrates the frequency dependence expected from good, flat samples.

The effects of wedge-shaped samples on the Cleveland method for determining resonance field values have been compared with the theoretical predictions and good qualitative agreement has been found.

C. Fermi Surface Calipers and Orbit Assignments

Fermi surface calipers have been determined from the raw RFSE angular data recorded and the thickness of the samples used. Where symmetry problems occurred in the angular data, complementary data have been recorded to allow systematic examination of both data sets to determine which features are common to both. These features have then been used to determine FS calipers resulting from RFSE.
The calipers have been examined using the computer predictions for second-zone calipers and the computer FS contour overlay plots to determine the origins in k-space of the various signals. The angular ranges and dependences for the experimental signals will be compared with those resulting from the NFE computer results.

The caliper data will be presented only on a rectangular coordinate system of caliper versus angle. Polar plots have not been used because for the second-zone hole surface of aluminum the caliper dimensions do not occur in a common plane. Thus the calipers do not correspond to FS dimensions in a single plane perpendicular to the normal. This problem in data interpretation has been discussed in more detail in an earlier section on the RPSE. As a result of this problem for aluminum the calipers have been plotted on rectangular coordinates to facilitate an easy comparison to the signal predictions resulting from the computer calculations.

Comparison of the experimental calipers will be made with the NFE second-zone hole surface. Recall that the NFE second-zone surface is less complicated than the actual second-zone FS. The difference in the two Fermi surfaces is the rounding of sharp edges on the actual surface resulting from interactions neglected in the NFE model. Estimates of rounding effects will be made and used to adjust the NFE calipers for further comparison with the experimental
Estimates of rounding can be made using the results of Pippard (42) and Kamm and Bohm (36). These studies yielded information on rounding on two different types of edges on the second-zone FS. First examine the work of Pippard and designate the edge considered as Type I. Figure 48a shows a second-zone cross section in a (100) plane through the center of the BZ. The corner examined by Pippard is located at the symmetry point W and is indicated in the figure. The rounding estimate made at this corner is also used for the edge of the second-zone FS between the symmetry points W and U. This rounding occurs at the intersection of a FS edge common to two hexagonal faces and a square FS face perpendicular to a [010] axis. Utilizing his information (42, Figure 11) a radius can be deduced for the rounding, $r=0.06\text{Å}^{-1}$. This leads to a reduction of $0.04\text{Å}^{-1}$ in the distance from the center of the BZ to the corner being examined. Edges between square and hexagonal faces on the second-zone FS will be referred to as Type I edges in the remaining discussion.

The remaining second-zone FS edges are common to two hexagonal faces and will be referred to as Type II edges. Data on a Type II edge can be taken from Kamm and Bohm (36, Figure 4). Their data relate to the rounding of a Type II FS edge where it intersects a square face perpendicular to a [010] axis.
Figure 48. Examples of FS rounding which modifies the NFE second zone FS.
Such an edge is labeled in Figure 48b which illustrates the intersection of the NFE second-zone FS with a (110) plane through the center of the BZ. The indicated rounding decreases the distance from the zone center to the FS edge by roughl 0.12Å⁻¹.

Figure 48c illustrates how the caliper reduction can depend on the orientation of the electron orbit on the FS. Orbit "1" is perpendicular to the rounded edge and the caliper reduction is minimized. Orbit "2" makes an acute angle with the rounded edge and the caliper reduction is larger than seen for orbit "1".

It should also be noted that electron orbits crossing Type I edges must be considered carefully to see if the caliper reduction is appropriate. Consider an electron following a real space trajectory with the shape seen in Figure 48a. If the sample normal is along a cubic axis the rounding of the Type I edge does almost nothing to reduce the caliper. But if the sample normal passes through the indicated Type I edge, then the entire rounding contributes to the reduction of the FS caliper.

The data will now be discussed for [100], [110], and [111] sample normals in that order. The calipers and orbit assignments will be discussed in detail and then summarized for each sample.
1. [100] sample normal

The angular data used were taken using sample Al 12. Due to the lack of any frequency dependence for this sample, the resonance value of the magnetic field to be used in Equation 9 has been taken at the low-field end of the resonance feature. The low-field end of the resonance occurs for the thickest part of the sample so the thickness used in Equation 9 was 180 microns as seen in Figure 41.

Figure 49a contains the calipers determined in this study. Figure 49b shows the predicted calipers for this sample for the NFE second-zone model.

From the NFE model caliper "A1" is seen to have the values $2.69 \times 10^{-1}$ at $0^\circ$ and $3.01 \times 10^{-1}$ at $20^\circ$. The signal results from complete orbits around the second-zone FS as indicated in Figure 50. These orbits cross two Type I edges in a manner that is not expected to significantly reduce the calipers. Experimental calipers for "A1" were $2.78 \times 10^{-1}$ at $0^\circ$ and $3.03 \times 10^{-1}$ at $20^\circ$. Beyond $20^\circ$ the experimental signal became too weak to measure. The loss of signal strength is expected as seen in Figure 49b for caliper "A1". Agreement is within 3% at $0^\circ$ and 1% at $20^\circ$.

Caliper "B1" can be examined at $45^\circ$ for the two figures in a similar manner. The NFE caliper is $2.79 \times 10^{-1}$. The hole orbits contributing to this signal are indicated and labeled in Figure 50. Notice that these orbits cross two Type II
Figure 49. Experimental and theoretical calipers determined for a [100] sample normal
Figure 50. Orbit assignments for experimental calipers for a [100] sample normal
edges. Thus the caliper for this orbit will be reduced by at least two times the value deduced from the data of Kamm and Bohm. If the plane of the orbit does not intersect the Type II edge at right angles, then the caliper will be even further reduced due to the geometry involved. This is based on the assumption that the rounding seen by Kamm and Bohm applies for the entire edge common to two hexagonal faces and not just where the edge meets the square PS face. Utilizing the Type II rounding reduces the expected caliper to 2.55Å⁻¹. The experimental value of 2.49Å⁻¹ agrees to within 2%.

It should be noted that the calipers "A1" and "B1" appear to intersect in the region of 20°-23°. Theoretically this is expected to occur at about 25°. The edge rounding just examined would shift this intersection to a smaller angle as can be seen from Figure 50. Scaling the change in location of the edges on Figure 50 indicates the peak should be shifted about 3° toward smaller angles.

Between 30° and 38° the signal associated with caliper "B1" broadens and is not distinct enough to allow a measurement. This is not anticipated from the computer predictions of signal intensity. The weakening may be related to the interaction of signals "B1" and "D1" in this angular range.

Caliper "C1" follows the same angular range on both the predicted and observed data. This caliper results from a signal that does not have an easily selected low-field edge.
This fact contributes to some of the scatter in this data. The angular dependence can be inferred from the next higher feature on the data trace which is also associated with this signal. Its minima is rather broad and occurs between 16° and 23°. This is compatible with the minima seen on Figure 24 which is broad and occurs between 15° and 22°. The electron orbit giving rise to caliper "C1" is indicated in Figure 50. It is seen that this orbit crosses one Type I edge in a direction that will contribute very little to a caliper reduction. It also crosses a Type II edge which will cause a reduction of about 0.12\(\text{Å}^{-1}\) in the expected caliper. Therefore, the NFE caliper at the minima is 2.31\(\text{Å}^{-1}\) and the adjusted NFE value is 2.19\(\text{Å}^{-1}\). The experimental value at the minima is 2.12\(\text{Å}^{-1}\) which agrees to within 3%.

Caliper "D1" on Figure 49a falls below its expected magnitude and also appears shifted to smaller angles than expected. The orbit thought to be contributing to this signal is shown in Figure 50. This orbit crosses the same edge combination as orbit "C1" so the rounding adjustments to the NFE caliper is expected to be 0.12\(\text{Å}^{-1}\). The resulting NFE and adjusted NFE calipers at the minima (27°) are 2.56\(\text{Å}^{-1}\) and 2.44\(\text{Å}^{-1}\) respectively. The experimental minima of 2.36 occurs at 24°. The agreement is within 3%.

No other signals have been identified in the complementing data sets that result from complete orbits.
around the second-zone hole surface. Calipers near $1.50\text{Å}^{-1}$ at $0^\circ$ in Figure 49b were not found but these signals are predicted to be much weaker than the signals already discussed.

Some signals have been attributed to partial orbits on the second-zone hole surface. Signals of this type have been discussed by Gantmakher (60) and have been predicted in aluminum by Druyvesteyn and Smets (123). They occur when an electron on an orbit moves parallel to the sample's surface at points other than extrema on the FS. If the electron orbit is such that it moves parallel to the surface at each surface, it can yield a RFSE signal even though it does not complete a full orbit. The important factor is for the particle's motion to remain near the surface long enough to interact with the applied fields. Examples of three partial orbits are shown in Figure 51. These regions on the FS can easily be selected in the computer FS contour overlay plots. The electron's velocity at a point on the FS is always perpendicular to the FS at the point, so velocities parallel to the surface of the sample will occur for FS regions where the FS is parallel to the normal axis. On the contour overlay plots the normal axis is perpendicular to the page. Thus regions where contour lines are very dense will correspond to the particle's velocity being almost parallel to the sample's surface. Such a situation exists in Figure 50 on the four sides of the square face centered on the [100] axis, so it is
Figure 51. Examples of partial orbits that can contribute to RPSE signals
possible to use the contour overlay plots to great advantage when evaluating possible contributions from partial orbits. Caliper "E1" on Figure 49a will be compared to the caliper resulting from the partial orbit "E1" on Figure 50. This orbit is qualitatively sketched in Figure 51c. From the NFE model in Figure 50 it is seen that caliper "E1" should contribute signals over the angular range of 190 to 450 and it crosses a Type I and a Type II edge at the limits of its real space motion. The Type I edge crossing is not expected to cause a caliper reduction due to the geometry involved. Notice that as the angle decreases toward 200, orbit "E1" begins to include motion across the square face near one of the sharp NFE corners. This corner is the same one studied by Kamm and Bohm so this caliper will be reduced by a second Type II correction in this angular region. The Type II edges will cause a caliper reduction as discussed earlier. The NFE calipers are 2.471-1 at 190 and 1.861-1 at 450. Including rounding effects reduces these to 2.251-1 and 1.751-1 respectively.

The experimental calipers designated "E1" cover the same angular range but their values fall below the expected values. They vary from 1.901-1 at 200 to 1.561-1 at 450. The experimental values are 15% and 10% too small at 200 and 450 respectively. The differences in magnitudes are not understood. Effects of rounding have already been included in
finding the expected values and the caliper differences are much too large to attribute them to further refinements in the rounding picture. If adjustments in the rounding model were made to account for this difference, the agreement for calipers already discussed would be destroyed. So no reasonable adjustments are seen and the differences remain unexplained. For this reason it is considered unlikely that this assignment is correct. But a careful examination of the FS has been made and another explanation has not been found for these data.

A low-field feature exists in the data sets presented in Figures 42 and 43 that also does not agree with a caliper that might be reasonably expected from the second zone. There is some disagreement between the complementing data sets for angles less than 10°. This feature shows the form expected for an orbit crossing the square face on the [100] axis but the caliper observed for 45° is about 30% smaller than expected from an examination of the contour overlay plot. Examination of the recorded data for this feature reveals that it does not have a line shape generally seen in the RFSE. In fact it appears to be associated with the background curve for \((df/dH)\) versus \(H\). The feature corresponds to the region where the slope of the background changes from positive to negative. For these reasons it is felt that these features are related to the processes governing the
background curve and not electron orbits spanning the sample's thickness. Therefore a caliper based on this feature is not shown.

Figures 42 and 43 also show several regions of high signal content for $\theta=45^\circ \pm 10^\circ$ and magnetic field values of 600 to 750 Oe and 900 to 1100 Oe. Some of these detailed variations may result from the small third-zone FS arms but the signals are too broad to allow an adequate treatment of the data to resolve the question.

The lack of symmetry seen in Figures 42 and 43 has made caliper assignments very difficult except for the strongest signals. The distortions placed on the normal four-fold symmetry have reduced the number of signals considered for caliper evaluation. It is very possible that several valid signals have been ignored because of their interaction with other signals not possessing the proper symmetry.

Further complications have been added resulting from samples that were not as flat as desired. The signal broadening caused by this has contributed to signal overlap in many regions and reduced the clarity of the angular dependence in such regions.

Caliper values and a comparison of the theoretical and experimental values are summarized in Table 4.
### Table 4. Caliper results for [100] sample normal

<table>
<thead>
<tr>
<th>Caliper</th>
<th>Angle (Degrees)</th>
<th>NFE caliper ((\bar{A}^{-1}))</th>
<th>Adjusted NFE caliper ((\bar{A}^{-1}))</th>
<th>Experimental caliper ((\bar{A}^{-1}))</th>
<th>Difference (((3)-(2)) \over (2))</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;A1&quot;</td>
<td>0</td>
<td>2.69</td>
<td>2.69</td>
<td>2.78</td>
<td>+3</td>
</tr>
<tr>
<td>&quot;A1&quot;</td>
<td>20</td>
<td>3.01</td>
<td>3.01</td>
<td>3.03</td>
<td>+1</td>
</tr>
<tr>
<td>&quot;B1&quot;</td>
<td>45</td>
<td>2.79</td>
<td>2.55</td>
<td>2.49</td>
<td>-2</td>
</tr>
<tr>
<td>&quot;C1&quot;</td>
<td>18</td>
<td>2.31</td>
<td>2.19</td>
<td>2.12</td>
<td>-3</td>
</tr>
<tr>
<td>&quot;D1&quot;</td>
<td>25</td>
<td>2.56</td>
<td>2.44</td>
<td>2.36</td>
<td>-3</td>
</tr>
<tr>
<td>&quot;E1&quot;</td>
<td>19</td>
<td>2.47</td>
<td>2.23</td>
<td>1.90</td>
<td>-15</td>
</tr>
<tr>
<td>&quot;E1&quot;</td>
<td>45</td>
<td>1.86</td>
<td>1.74</td>
<td>1.56</td>
<td>-10</td>
</tr>
</tbody>
</table>

2. **[110] sample normal**

FS calipers were determined using the angular data recorded with sample A1 7. The signals observed were very weak. For this reason the data do not represent all the signals expected for this sample orientation. The data seen have been related to expected signals as seen from the computer predictions for a [110] sample normal.

Due to the weakness of the signals it was very difficult to determine the low-field edge of the signals. For this reason the resulting FS calipers may be slightly larger than
the actual values. The error could be on the order of 0.10 Å⁻¹. This would cause a shift to smaller calipers but the angular dependence of the signals relative to each other would not be altered.

The caliper magnitudes have been calculated using Equation 9. The sample thickness taken from Figure 41c is 196 microns. The isolated reading of 198 microns has been neglected.

The angular variations of the calipers are shown in Figure 52a. They will be compared with the computer predictions for calipers of the second-zone hole surface as seen earlier in Figures 25 and 28.

Caliper "A2" from Figure 52a corresponds to the upturned signal shape seen at 0° and about 2.1Å⁻¹ on Figure 52b. This signal is attributed to the hole orbit indicated in Figure 53. The orbit crosses two Type I edges in a direction that should result in corrections close to the full estimated value of 0.04Å⁻¹ for each edge. The NPE and adjusted NPE calipers are then 2.09Å⁻¹ and 2.01Å⁻¹ respectively at 0°. The experimental caliper is 2.08Å⁻¹ and agrees with the adjusted NPE value within 3%.

From Figure 52b it is seen that a smooth extension of this signal intersects a higher caliper signal, "B2", at about 13°. This is compatible with a smooth extrapolation of the calipers "A2" and "B2" in Figure 52a.
Figure 52. Experimental and theoretical calipers determined for a [110] sample normal
Figure 53. Orbit assignments for experimental calipers for a [110] sample normal
Next examine caliper "B2". From Figure 52b it is seen that the predicted intensity of this signal is fairly low until an angle of about 20° is reached. Signal "B2" from the experimental data was not observed for angles less than 24°. From Figure 53 it is seen that both "B2" orbits cross one Type I edge and one Type II edge. The Type I edge is the same as discussed above and a caliper reduction of 0.04Å⁻¹ is expected. The Type II edge rounding is not very effective at reducing the caliper due to the orbit geometry. The estimated reduction of 0.05Å⁻¹ is based on a graphical examination of the edge under consideration. The net correction is then about 0.09Å⁻¹.

The NPE and adjusted NPE calipers for "B2" at 24° are 2.34Å⁻¹ and 2.25Å⁻¹ respectively. The experimental value of 2.36Å⁻¹ agrees to within 5%. A similar examination of NFE, adjusted NFE, and experimental values at the minima for "B2" yield 2.25Å⁻¹, 2.16Å⁻¹, and 2.27Å⁻¹ respectively. The agreement is again within 5%. Both the experimental and theoretical minima for "B2" occur at 34° ±1°. Based on the NPE second zone it is seen that caliper "B2" rises to 2.65Å⁻¹ near 55°. It then remains constant between 55° and 63°. The adjusted caliper value is about 2.56Å⁻¹. The experimental data shows signs of flattening out between 55° and 61° but the curve never does become flat. The value at 60° is 2.83Å⁻¹. The values differ by 11%.
An examination of caliper "B2" on Figure 52b shows that a lower branch splits away from "B2" as \( \theta \) increases beyond 43°. This signal, "C2", is then seen to extend upward toward 3.13Å\(^{-1}\) at 72°. Orbit "C2" crosses two Type II edges in the same manner just considered for "B2". Thus the rounding is expected to reduce the caliper by 0.10Å\(^{-1}\) to 3.03Å\(^{-1}\) at 72°. Such a signal was observed experimentally. The caliper "C2" extends to roughly 74° at 3.15Å\(^{-1}\) and agreement is within 4%. The hole orbit yielding signal "C2" is labeled on Figure 53.

A signal for the caliper "D2" on Figure 52a was observed but an angular study was not made in this field range. This feature corresponds to the expected signal "D2" seen on Figure 52b near 0° and 1Å\(^{-1}\) and the hole orbit is indicated in Figure 53. The orbit crosses two Type II edges and the reduction in caliper is expected to be the same as found for orbit "C2", 0.10Å\(^{-1}\). The NFE and adjusted NFE calipers are then 1.00Å\(^{-1}\) and 0.90Å\(^{-1}\) respectively. The experimental value of 0.95Å\(^{-1}\) agrees to within 4% with the adjusted NFE value.

Signals "E2" and "F2" on Figure 52b were not observed even though they are expected to be fairly strong. This is a direct result of the experimental sensitivity for the RPSE being zero for 90° in Figures 52a and 52b.

Recall that due to the weakness of the signals it was extremely difficult to determine the low-field feature of the
signals. For this reason the experimental values for the calipers should probably be slightly smaller than those indicated above.

The agreement seen in the angular behavior is of importance. The comparison of the expected and observed calipers showed maxima, minima, and signal splittings at the proper angular locations. All the signals seen for this sample agreed with predicted calipers and were due to complete hole orbits around the second-zone hole surface. No signals were seen that have not been discussed.

Due to the weakness of the signals an attempt was not made to rotate the sample in the coil and take data on the region near 90° in Figures 52a and 52b.

Caliper values and a comparison of the theoretical and experimental values are summarized in Table 5.

3. [111] sample normal

Several [111] samples were used to obtain data. Signals were observed whose field values varied with sample thickness as required by Equation 9. The data to be discussed were recorded using Al 16. This sample yielded the largest number of signals and the best signal strength of the samples examined. Again complementary data sets (Figures 44 and 45) have been used in an effort to select useful data and to eliminate data whose symmetry does not agree with the FS symmetry. Calipers have been calculated using a sample
Table 5. Caliper results for [110] sample normal

<table>
<thead>
<tr>
<th>Caliper Angle (Degrees)</th>
<th>NPE caliper ( \langle \tilde{A} \rangle^{-1} )</th>
<th>Adjusted NPE ( \langle \tilde{A} \rangle^{-1} )</th>
<th>Experimental caliper ( \langle \tilde{A} \rangle^{-1} )</th>
<th>Difference ( \frac{(3) - (2)}{(2)} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;A2&quot; 0.</td>
<td>2.09</td>
<td>2.01</td>
<td>2.08</td>
<td>+3</td>
</tr>
<tr>
<td>&quot;B2&quot; 24.</td>
<td>2.34</td>
<td>2.25</td>
<td>2.36</td>
<td>+5</td>
</tr>
<tr>
<td>&quot;B2&quot; 34.</td>
<td>2.25</td>
<td>2.16</td>
<td>2.27</td>
<td>+5</td>
</tr>
<tr>
<td>&quot;C2&quot; 72.</td>
<td>3.13</td>
<td>3.03</td>
<td>3.15</td>
<td>+4</td>
</tr>
<tr>
<td>&quot;D2&quot; 0.</td>
<td>1.00</td>
<td>0.90</td>
<td>0.95</td>
<td>+6</td>
</tr>
</tbody>
</table>

thickness of 120 microns as taken from Figure 41d.

Caliper "A3" on Figure 54a has been attributed to holes completing motion across a hexagonal second-zone PS face perpendicular to the normal axis. This partial orbit is illustrated and labeled as "A3" in Figure 55 and is similar to that sketched in Figure 51b. The caliper varies smoothly over the angular range from 6° to 30°.

The orbit crosses one Type II edge but the geometry involved for this sample decreases the caliper reduction due to rounding on this edge. From a graphical examination of a Type II edge for this orientation, the caliper reduction is estimated to be about 0.02\( \tilde{A}^{-1} \). It also crosses one Type I
Figure 54. Experimental and theoretical calipers determined for a [111] sample normal
Figure 55. Orbit assignments for experimental calipers for a [111] sample normal.
edge but the geometry involved yields no caliper reduction for the rounding of this edge. The resulting net reduction to the caliper is estimated to be 0.02Å⁻¹. The NFE and adjusted NFE calipers at 30° are then 1.65Å⁻¹ and 1.63Å⁻¹ respectively. The experimental value of 1.59Å⁻¹ is in agreement to within 2%.

The data also have been compared with the adjusted NFE calipers for this signal at 10° because both curves are still smoothly varying at 10°. The NFE and adjusted NFE calipers are 1.82Å⁻¹ and 1.80Å⁻¹ while the experimental value is 1.72Å⁻¹. Agreement is within 4%.

For this angular range an inspection of Figure 55 shows that orbits of type "A3" cross opposing edges of the FS that are almost parallel to each other. Thus many orbits have nearly the same caliper and the signal strength is enhanced.

An examination of the data traces indicates that signal "A3" can be positively identified over the region from 6° to 30°. For less than 6° the feature broadens and identification is no longer possible. This signal loss could be a direct result of decreasing the number of holes contributing to the signal as the number of orbits of type "A3" crossing opposite edges of the hexagonal FS face.

Caliper "B3" in Figure 54a is also due to a partial orbit crossing the hexagonal face of the second-zone FS which is perpendicular to the sample's normal. The orbit crosses
the hexagonal face and one square face of the second-zone hole surface. It is illustrated in Figure 55. The caliper increases smoothly as the angle changes from $30^\circ$ toward $6^\circ$ where caliper maxima occurs. Examine caliper "B3" in Figure 55. The caliper is seen to reach a maxima at about $6^\circ$ as the orbit moves over the point of the PS located between the $[\bar{2}11]$ axis and the $[1\bar{1}0]$ axis. Between $6^\circ$ and $30^\circ$ the orbit crosses a Type II edge in the same geometry discussed for orbit "A3", so the resulting caliper reduction is estimated to be $0.02\AA^{-1}$. It also crosses a Type I edge in a manner that will yield a maximum caliper reduction for that type edge, $0.04\AA^{-1}$. The net reduction is then $0.06\AA^{-1}$, but for about $6^\circ$ the Type I edge reduction is complicated by a merger with a Type II edge. This corresponds to the rounding encountered at the corner of a square face where a Type II edge intersects two Type I edges. From a graphical method the rounding of this corner as seen at $6^\circ$ is expected to be $0.08\AA^{-1}$ and the net reduction will be $0.10\AA^{-1}$.

At $30^\circ$ the NPE and adjusted NFE calipers are $2.20\AA^{-1}$ and $2.14\AA^{-1}$ respectively. The corresponding experimental value is $2.16\AA^{-1}$ resulting in agreement within 1%. Similarly at $6^\circ$ the expected values are $2.68\AA^{-1}$ and $2.58\AA^{-1}$. The experimental value is $2.50\AA^{-1}$ and agreement is within 3%.

For angles between $6^\circ$ and $0^\circ$ the orbit crosses the same Type II edge already considered (a reduction of $0.02\AA^{-1}$) and
another Type II edge that should give almost its full reduction of \(0.12 \AA^{-1}\). The resulting NFE and adjusted NFE calipers at \(0^\circ\) are \(2.46 \AA^{-1}\) and \(2.32 \AA^{-1}\) respectively. The experimental value of \(2.42 \AA^{-1}\) is taken at \(2.5^\circ\) due to a lack of signal strength at \(0^\circ\). Agreement is within 4%. The experimental signal washed out near \(0^\circ\) and did not result in a caliper value but the caliper is not expected to change significantly in the \(2.5^\circ\) interval.

Calipers "C3" and "D3" in Figure 54a have been attributed to complete orbits and are expected from the predicted signals of Figure 54b. The orbits contributing to these calipers are indicated on the FS contour overlay plot of Figure 55. These orbits are of special interest due to the fact that Gantmakher and Krylov (89) did not see similar orbits for their indium sample with a \([111]\) normal axis. They indicated (89, Figure 7) no calipers resulting from complete orbits for that sample. Orbits similar to "C3" and "D3" should exist for indium as the second-zone hole surface is very similar to that for aluminum. The major difference is the distortion of the indium BZ as a result of its tetragonal crystal structure.

Caliper "C3" is predicted for angles from \(0^\circ\) to about \(25^\circ\) (where the intensity decreases) with a minima at roughly \(18^\circ\). This caliper will be smaller than predicted due to rounding effects, but the reduction will vary with the direc-
tion of the orbits. For 0° it crosses two Type I edges and should be reduced by roughly 0.08Å⁻¹. Between 15° and 25° the orbit crosses one Type I edge and one Type II edge so the calipers should be reduced by about 0.16Å⁻¹. The resulting NPE and adjusted NPE calipers at 1.3° are 3.03Å⁻¹ and 2.95Å⁻¹ respectively. The experimental value of 2.91Å⁻¹ agrees to within 1%. The comparison was made at 1.3° due to the lack of experimental data at 0°. At 20° the NPE and adjusted NPE calipers are 2.59Å⁻¹ and 2.43Å⁻¹ respectively. The experimental value is 2.50Å⁻¹ which agrees within 3%.

From Figure 54b it is seen that caliper "D3" is expected to occur for the angular range of 15° to 30° but the signals are strongest for angles greater than 20°. This orbit crosses two square faces of the second zone FS during its entire range so the caliper reduction is determined by two Type I edges. The geometry involved leads to the expected reduction for each edge being the maximum Type I reduction discussed earlier, 0.04Å⁻¹. Therefore, the net reduction is estimated to be 0.08Å⁻¹. The NPE and adjusted NPE at 20° are then 2.87Å⁻¹ and 2.79Å⁻¹ respectively. The experimental value of 2.83Å⁻¹ agrees to within 1%. At 30° the three caliper values are 2.78Å⁻¹, 2.70Å⁻¹, and 2.70Å⁻¹ respectively.

Caliper "E3" is attributed to a partial orbit across a square face of the second-zone FS. The angular dependence of
the caliper is shown in Figure 54a and the corresponding hole orbit is indicated on Figure 55. The holes contributing to this signal cross two Type I edges but the geometry of this case is expected to yield no caliper reduction due to this rounding. The only caliper reduction is expected to occur near 30° where the effects of the Type II edge rounding will round the corners of the square face itself. From a graphical examination this is not expected to affect calipers for angles smaller than 25°, so for the angular range of 0° to 24° there are no corrections to the NFE calipers. At 0° the NFE and experimental calipers are 0.92Å⁻¹ and 0.86Å⁻¹ respectively. Agreement is within 7%. At 24° the NFE and experimental values are 1.05Å⁻¹ and 0.98Å⁻¹ and agreement is again within 7%. The differences seen here in the expected and experimental values can not be removed by simply adjusting the edge rounding. In the geometry involved in this specific case, the calipers are very insensitive to the effects of rounding the edges. The intensity of the signal is expected to decrease near 25° as the number of electrons that can contribute to the signal decreases as the orbits near the corners on the square face. A corresponding loss of signal is seen experimentally. This caliper assignment appears to be the only one possible after a thorough examination of the second-zone surface. But the insensitivity of this orbit to rounding effects and the 7% difference between
observed and expected calipers make it seem unlikely that the assignment is correct.

One additional signal occurs that has not been associated with a hole orbit on the second-zone FS. This signal is seen on Figures 44 and 45 near 1100 Oe at 30° ±15°. The feature reaches a peak at roughly 1100 Oe and 30°. This corresponds to caliper of 1.96Å^-1 after field corrections have been included. A second-zone orbit has not been identified that would be compatible with the field values or the angular dependence seen in this signal. It is felt that the signal is probably due to a coupling of a third-zone arm orbit and the partial orbit "A3" seen on Figures 54a and 55. The arm diameter would then be compatible with estimates made from the de Haas-van Alphen data of Larson and Gordon (19). The width of the signals for this sample again makes it very difficult to isolate signals due to different electron orbits. The difficulty in isolating signals from separate orbits in regions like this decreases the chance to resolve regions of this type.

The [111] sample normal calipers are summarized in Table 6.

4. Discussion

A comparison of the calipers obtained in this investigation with previous data is not simply a matter of numerical comparison. As discussed earlier, the RFSE caliper
Table 6. Caliper results for [111] sample normal

<table>
<thead>
<tr>
<th>Caliper</th>
<th>Angle (Degrees)</th>
<th>NFE caliper (Å⁻¹)</th>
<th>Adjusted NFE caliper (Å⁻¹)</th>
<th>Experimental NFE caliper (Å⁻¹)</th>
<th>Difference (Å⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;A3&quot;</td>
<td>0.</td>
<td>2.13</td>
<td>2.05</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>&quot;A3&quot;</td>
<td>10.</td>
<td>1.82</td>
<td>1.80</td>
<td>1.72</td>
<td>-4</td>
</tr>
<tr>
<td>&quot;A3&quot;</td>
<td>30.</td>
<td>1.65</td>
<td>1.63</td>
<td>1.59</td>
<td>-2</td>
</tr>
<tr>
<td>&quot;B3&quot;</td>
<td>6.</td>
<td>2.68</td>
<td>2.62</td>
<td>2.50</td>
<td>-3</td>
</tr>
<tr>
<td>&quot;B3&quot;</td>
<td>30.</td>
<td>2.20</td>
<td>2.14</td>
<td>2.16</td>
<td>+1</td>
</tr>
<tr>
<td>&quot;C3&quot;</td>
<td>1.3</td>
<td>3.03</td>
<td>2.95</td>
<td>2.91</td>
<td>-1</td>
</tr>
<tr>
<td>&quot;C3&quot;</td>
<td>20.</td>
<td>2.59</td>
<td>2.43</td>
<td>2.50</td>
<td>+3</td>
</tr>
<tr>
<td>&quot;D3&quot;</td>
<td>20.</td>
<td>2.87</td>
<td>2.79</td>
<td>2.83</td>
<td>+1</td>
</tr>
<tr>
<td>&quot;E3&quot;</td>
<td>0.</td>
<td>0.92</td>
<td>0.92</td>
<td>0.86</td>
<td>-7</td>
</tr>
<tr>
<td>&quot;E3&quot;</td>
<td>24.</td>
<td>1.05</td>
<td>1.05</td>
<td>0.98</td>
<td>-7</td>
</tr>
</tbody>
</table>

corresponds to the distances across the shadow of the FS projected on a plane perpendicular to the sample's normal axis. For the second-zone FS of aluminum this means that many calipers do not correspond to a distance across the FS in a single plane perpendicular to the normal axis. Consequently many of the calipers do not correspond to FS dimensions in a plane passing through the center of the BZ.
For this reason the calipers have been compared with calipers derived from the NPE model for the second-zone hole surface. This model has been carefully examined in other studies listed earlier with the only resulting modification being the rounding of sharp edges on the FS. The edge rounding indicated by the studies of Kamm and Bohm (36) and Pippard (42) have been utilized when comparisons have been made with the NPE FS model. These rounding estimates were based on the assumption that the rounding of an edge remained the same at every point on that edge. This assumption allowed a phenomenological adjustment to the NPE second-zone calipers.

Before examining the agreement of the present data with the adjusted NPE model for the second-zone FS, it is important to examine possible sources of experimental error in this study.

The accuracy of the experimental FS calipers given in Tables 4, 5, and 6 are limited by the accuracy of the values of magnetic field and sample thickness used in Equation 9.

The accuracy of the magnetic field values was subject to two factors. First, for the range of field values used in this study the magnet supply accuracy was limited to ±10 Oe. This 10 Oe uncertainty translates into a caliper uncertainty of 0.02Å⁻¹ to 0.03Å⁻¹ for the thickness ranging from 120 to 196 microns. Thus for the calipers shown in Tables 4, 5, and
this uncertainty ranges from less than 1% to greater than 3% depending on the sample thickness and the magnitude of the observed calipers.

The second factor of magnetic field uncertainty was related to the accuracy of the selected value of the magnetic field associated with the onset of the RFSE resonance. When visually selecting the point on a curve where it deviates from the background curve, a finite amount of deviation is needed for detection. This tends to cause field values to be too large. This could contribute to a systematic error in the caliper values by making them too large. This factor was small for sharp features on a RFSE data trace but relatively large for very broad signals. It has already been noted that the data recorded for the [110] sample normal were very weak and it was not possible to determine the onset of the RFSE signal. Instead, the first extrema of the resonance was measured for calipers "A2", "B2", and "C2". From the shapes of the data traces it is felt that the onset of the resonance would correspond to calipers smaller than those given in Table 5 by 0.09\textup{100}^{-1} to 0.12\textup{100}^{-1}. This correction would decrease the differences seen between the experimental and adjusted NPE caliper values.

Signals used to determine the calipers for [100] and [111] normal samples were much stronger and this problem was not of major importance. These (df/dH) versus H traces were
recorded twice for each angle to check the reliability of the data and the resulting uncertainty in the calipers is felt to be less than ±0.02Å⁻¹ due to mismeasurement of signal features.

The value of the sample thickness used in Equation 9 resulted from a series of values recorded on a square grid pattern for each sample. For a sample with non-uniform thickness it is very unlikely that a measurement point would coincide with the thickest spot of the sample. In regions not adjacent to the sample edges the changes in thickness occur slowly so it is felt that this could change the thickness by 2 microns. Such a difference would increase the calculated calipers by 1.1% for the [100] sample, 1.0% for the [110] sample, and 1.7% for the [111] sample. With this must be included the uncertainty of the recorded values taken with the microscope of ±2 microns. The net effect of these uncertainties on the caliper error bars is for the [100] sample -1.1% to +2.2%, for the [110] sample -1.0% to +2.0%, and for the [111] sample -1.7% to +3.4%.

It should also be noted that the NFE calipers as contained in Figures 24, 25, and 26 introduce some uncertainty as the values have been grouped in intervals that are 0.03125Å⁻¹ wide. Thus the caliper value assumed for a given interval may differ from the actual value by ±0.03Å⁻¹. This must be considered when the agreement between the adjusted
NFE and the experimental calipers is examined.

The experimental calipers as seen in Figures 49a, 52a, and 54a exhibit very little scatter from point to point. The uncertainties mentioned above are thus related to the calibration of all the caliper values and not the uncertainty in the calipers as seen from point to point (i.e. systematic errors).

The experimental and adjusted NFE second-zone calipers are compatible when error bars are evaluated for the two sets of calipers. For the [110] normal sample it was also necessary to estimate the caliper reduction associated with finding the onset of the BFSE signal. The FS edge rounding estimates used have lead to consistent agreement between the two sets of calipers. The current data indicate that no changes in the adjusted NFE second-zone FS need be considered to obtain the observed angular dependence and magnitude for the PS calipers.
VII. CONCLUSIONS

The Fermi surface of aluminum has been studied by the RFSE technique using samples with normal axes along the three major symmetry axes. The resulting FS calipers have been examined and many have been identified with specific hole orbits on the second-zone hole surface. The caliper magnitudes and angular dependence observed are in good agreement with the NFE second-zone FS when FS edge rounding is taken into account.

In the course of this investigation the use of frequency studies was examined as a useful tool to determine the resonant field values for recorded data. The method employed by Cleveland (51) was examined in detail and found to yield the proper results only for perfectly flat samples. Non-uniformity in sample thickness was found to cause the resulting field value and exponent of frequency dependence to be artificially low. This was experimentally substantiated.

The effects of non-uniform sample thickness were also observed on the signals recorded in the RFSE. It was seen that a wedge-shaped sample resulted in signal splitting while samples with a non-uniform but random thickness give rise to signal broadening and signal distortion.
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