Secondary electron emission in the scanning electron microscope\textsuperscript{a)}

H. Seiler

Institut für Physik der Universität Hohenheim, D-7000 Stuttgart-70, Garbenstr. 30, West Germany

(Received 7 January 1983; accepted for publication 15 July 1983)

This paper surveys experimental results concerning secondary electron emission of surfaces bombarded by primary electrons with respect to scanning electron microscopy. The energy distribution, the angular distribution, and the yield of secondary electrons from metals and insulators are reviewed as well as the escape depth of the secondary electrons and the contribution of the backscattered electrons to the secondary electron yield. The different detectors for secondary electrons in the scanning electron microscope are described. The contrast mechanisms in the scanning electron microscope, material, topography, voltage, magnetic, and crystallographic orientation contrast based on secondary electron emission, as well as the lateral resolution, depending among other things on the spatial distribution of the emitted secondary electrons, are discussed.

PACS numbers: 79.20.Hx, 07.80. + x

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INTRODUCTION

For imaging of surfaces in the scanning electron microscope (SEM) mostly the “true” or slow secondary electrons (SE) are used. These are electrons which leave the surface with energies $E_{SE} < 50$ eV when the object is bombarded with primary electrons (PE). The SE yield depends on the material and the topography of the specimen. The SE can be de-

\textsuperscript{a) Dedicated to Professor Dr. Gerhard Pfefferkorn on the occasion on his 70th birthday.}

![Diagram of SE emission](image)

FIG. 1. Arrangement for measuring the SE emission. The secondary electrons (SE) are released by primary electrons (PE) and by backscattered or reflected electrons (RE). The mean escape depth $\lambda$ of the SE is about $\lambda \approx 1$ nm from metals. With the detector $D$ all emitted electrons or only the RE can be measured.
by several methods. The specimen is bombarded with primary electrons (PE). All emitted electrons independent of their emission direction are collected. It is possible to separate the slow from the energetic backscattered or reflected electrons (RE) by a retarding field. The grid is held negative with respect to the collector so that secondaries formed at the collector do not disturb the measurements. The slow SE are not only released by PE at the entry point but also by RE at the exit point from the specimen.

Figure 2 shows the specimen chamber of a SEM. The sample has a rough surface and underlying object details (such as A and B in Fig. 2) of different material or different crystallographic grains provide image contrasts. Although the escape depth of the SE is very small, object details deeper than the escape depth influence the SE emission and are visible in SEM because the RE also excite SE. The object is normally not in UHV and absorbed layers may be at the surface. So by bombarding the specimen with a PE beam with a high current density contamination or desorption of the absorbed layers may occur. Both effects may influence the SE emission depending on bombarding time. The SE are extracted by electric fields to one or several detectors. The image contrast from rough surfaces can be improved if the strength of the electric field is insufficient to collect emitted SE. The signal at the detector contains different contributions as shown in Fig. 2.


I. EXPERIMENTAL DATA ON SE EMISSION

If a surface is bombarded with primary electrons (PE), electrons are released. Figure 3 shows schematically the energy distribution of these electrons released by PE with energies 100 eV < $E_{PE}$ < 1 keV. With increasing energy of the PE the elastic peak decreases and a broader maximum of inelastic PE is to be seen. According to their energy the electrons can be divided in different groups:

1. Electrons with energies $E < 50$ eV: Secondary electrons (SE);
2. Electrons with energies 50 eV < $E < E_{RE}$: Inelastically backscattered and elastically reflected electrons (RE).

According to the different groups we can define

1. SE yield $\delta$: number of SE/number of PE;
2. Backscattering coefficient $\eta$: number of RE/number of PE;
3. Total electron yield $\sigma = \delta + \eta$;
4. Coefficient of elastically reflected electrons $\eta_{E}$: number of elastically reflected electrons/number of PE (Schmid et al. 1983).

Superimposed on the energy distribution there are often some peaks. Some of these are Auger electrons (having an

FIG. 3. Schematic energy distribution of electrons which are emitted from a surface bombarded with PE. The SE yield is up to 10$^4$ times higher than the Auger electron yield.

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energy depending on the surface material) and others are electrons which are backscattered with an energy loss by volume or surface plasmon excitation or ionization of surface atoms.

A. Energy distribution of SE

The energy distribution of SE, released by PE with energies more than 100 eV is essentially independent of the energy of the PE, and is characterized by the most probable energy $E_{\text{SE}}^m$ and the full width at half maximum (HW). Ideally $E_{\text{SE}}^m$ should be measured relative to the Fermi energy level. Both $E_{\text{SE}}^m$ and HW depend on the surface material. HW is smaller for insulators than for metals and depends strongly on very thin surface layers (Dietrich and Seiler 1960). According to Kollath (1956), for metals one gets $1.0 \text{ eV} < E_{\text{SE}}^m < 2.5 \text{ eV}$ and $4 \text{ eV} < \text{HW} < 7 \text{ eV}$. New measurements on clean metal surfaces show $1 \text{ eV} < E_{\text{SE}}^m < 5 \text{ eV}$ and $3 \text{ eV} < \text{HW} < 15 \text{ eV}$ (Schäfer and Höflz 1972). Figure 4 shows typical energy distributions for metals and insulators. (Bouchard and Carette 1980).

According to Chung and Everhart (1974) the shape of the energy distribution of SE is given by

$$\frac{dN}{dE} = k(E - E_F - \Phi)/(E - E_F)^3,$$

where $k$ is the material constant, $E_F$ is the Fermi energy, and $\Phi$ is the work function.

B. Angular distribution of SE

The angular distribution (see Fig. 5) of SE from polycrystalline surfaces is a cosine distribution independent of the angle of incidence of the PE (Jonker 1957; Kanaya and Kawakatsu 1972). The angular distribution of SE of a single crystal face shows anisotropy (Burns 1960; Begrambekov et al. 1971) and the energy-angle distribution shows a sharp fine structure (Appelt 1968).  

C. SE yield of metals and insulators

Figure 6 shows schematically the SE yield as a function of primary energy $E_{\text{PE}}$. The general shape is the same for all materials: $\delta$ increases with $E_{\text{PE}}$, reaches a maximum value $\delta^m$ at $E_{\text{PE}}^m$, and then decreases with increasing $E_{\text{PE}}$. Table I shows values for $\delta^m$ and $E_{\text{PE}}^m$ at normal incidence (Seiler 1967, 1968a; Kanaya and Kawakatsu 1972; Kanaya et al. 1978; Ono and Kanaya 1979). For the values of the backscattering coefficient $\eta$ at $E_{\text{PE}}^m$ see Seiler (1967). For $E_{\text{PE}} > 10 \text{ keV}$, $\eta$ is constant and increases monotonically with the atomic number $Z$ (Ono and Kanaya 1979). This value is shown in Table I. The values $\delta$ (20 keV) were calculated using the semiempirical theory of SE with formula (16). The values are too high compared with experimental values. According to the measurements of Reimer et al. (1968), Seiler (1968a), we get for Al: $\delta$ (20 keV) 0.1–0.17 and for Cu: $\delta$ (20 keV): 0.14. However there are only few measurements available for PE energies used in SEM, so the calculated values are given in Table I.

![FIG. 5. Angular distribution of SE emitted from a polycrystalline surface.](image)

Some care must be taken comparing the values for $\delta^m$ of different authors because often the maximum total yield $\sigma^m$ is given instead $\delta^m$. The values for $\delta^m$ are about 0.35 and 1.6 for metals at 100 eV $< E_{\text{PE}} < 800$ eV and between 1.0 and 10 for insulators at 30 eV $< E_{\text{PE}} < 2000$ eV. High values for $\delta$ are found on single crystals of insulators such as MgO: $\delta^m \approx 20–25$. $\delta^m$ reaches high values if $E_{\text{PE}}^m$ is also high. For metals we get $\delta^m/E_{\text{PE}}^m \approx 2 \times 10^{-3}$ eV (Ono and Kanaya 1979). An example for the simultaneous increase of $\delta^m$ and $E_{\text{PE}}^m$ was shown by Beisswenger and Gruner (1974) at reactive evaporated BeO layers. There exists no simple relation

![FIG. 6. SE yield $\delta$ depending on $E_{\text{PE}}$.](image)
between $\delta$ and the atomic number $Z$ of the surface atoms, so material analysis by measuring the SE yield is not possible. Many authors tried to show relations between $\delta^m$ and regularities of the elements of the periodic table (e.g., Seiler 1967; Ono and Kanaya 1979; Makarov and Petrov 1981). Materials consisting of atoms with a large diameter have small $\delta^m$ (Seiler 1967). According to Ono and Kanaya (1979), $\delta^m$ and $E_{PE}$ depend both on the ionization energy $J$ of the surface atoms: $\delta^m \sim J^{4/5}$, $E_{PE} \sim J^{4/3}$.

A survey on some measured values of $E_t < E_{PE}$ and $E_{II} > E_{PE}$ for which $\delta = 1$ at some metal surfaces are given by Whetten (1961):

- Ag: $E_t = 200 \text{ eV}$, $E_{II} > 2000 \text{ eV}$
- Au: $E_t = 300 \text{ eV}$, $E_{II} > 2000 \text{ eV}$
- Pt: $E_t = 350 \text{ eV}$, $E_{II} = 3000 \text{ eV}$
- Si: $E_t = 125 \text{ eV}$, $E_{II} = 500 \text{ eV}$
- Cu: $E_t = 200 \text{ eV}$, $E_{II} = 1500 \text{ eV}$

Table II shows some values of $\sigma^m$ and $E_{PE}$ for compounds and insulators. Measurements of the SEE of insulators are difficult to make because of charging effects. For the investigation of insulators and electrical floating objects the values of $E_{PE}$ where $\sigma = 1$ are important (see $E_{PE}$ and $E_{II}$ in Fig. 6). Using these PE energies the potential of the surface will stabilize. For $E_{PE} > E_{II}$, as normally used in the SEM, the potential of the surface becomes negative because $\sigma < 1$; more electrons reach the surface than SE and RE leave the surface. The PE are decelerated, thus increasing $\sigma$ until $\sigma = 1$. The negative potential of the surface can reach very high values up to the potential of the electron gun. For $E_{PE} < E_{II}$, the sample becomes positively charged because $\sigma > 1$. The PE are accelerated until $\sigma = 1$ is reached and so the crossover for $E_{II}$ gives a stable point. The positive potential however can only reach values of the collector potential, otherwise the SE will return to the surface.

The SE yield of thin insulating layers on a conducting bulk material increases by positive charging of the surface. If

<table>
<thead>
<tr>
<th>Atom</th>
<th>$\delta^m$ (eV)</th>
<th>$E_{II}$ (eV)</th>
<th>$E_{PE}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>CuO</td>
<td>1.2</td>
<td>400</td>
<td></td>
</tr>
<tr>
<td>PbS</td>
<td>1.2</td>
<td>500</td>
<td></td>
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<td>1.0-1.2</td>
<td>500</td>
<td></td>
</tr>
<tr>
<td>MoO3</td>
<td>1.1-1.3</td>
<td>450</td>
<td></td>
</tr>
<tr>
<td>CsCl</td>
<td>1.5</td>
<td>500</td>
<td></td>
</tr>
<tr>
<td>Al2O3</td>
<td>2.6-4.7</td>
<td>600</td>
<td></td>
</tr>
<tr>
<td>SiO2</td>
<td>2.1-2.9</td>
<td>400</td>
<td></td>
</tr>
<tr>
<td>ZnS</td>
<td>1.8</td>
<td>350</td>
<td></td>
</tr>
<tr>
<td>Mica</td>
<td>2.4</td>
<td>350</td>
<td></td>
</tr>
<tr>
<td>Glass</td>
<td>2-3</td>
<td>300-450</td>
<td></td>
</tr>
<tr>
<td>BaF2</td>
<td>4.5</td>
<td>500</td>
<td></td>
</tr>
<tr>
<td>SrO</td>
<td>1.5</td>
<td>700</td>
<td></td>
</tr>
<tr>
<td>GeO</td>
<td>3.4-10</td>
<td>400-2000</td>
<td></td>
</tr>
<tr>
<td>BaO</td>
<td>2.3-4.8</td>
<td>400</td>
<td></td>
</tr>
<tr>
<td>CaO</td>
<td>2.2</td>
<td>500</td>
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<tr>
<td>KBr</td>
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</tr>
<tr>
<td>KCl</td>
<td>12</td>
<td>1600</td>
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</tr>
</tbody>
</table>

### Table II

- $\sigma^m$ and $E_{PE}$ for total SE yield of some compounds and insulators according to Whetten (1961).

### Table I

- $\delta^m$ maximum of the SE yield for different atomic numbers $Z$ and calculated values for $\delta$ and $\eta$ for typical energies used in the SEM.
FIG. 7. Decrease of $\delta$ of an aluminum surface at $E_{PE} = 20$ keV depending on the thickness of a contamination layer grown under electron bombardment.

the electric field strength within the layer exceeds $10^7$ V/cm the Malter (1936) effect sets in, i.e., the electron emission is caused by field emission at the bulk material through the insulating layer. In this case the electron yield becomes independent of the PE current. In the case of the field enhanced SEE the electric field is less than $10^7$ V/cm, so no field emission can occur—however, the escape possibility of the SE is increased. Seiler (1967) estimated that on thin insulating layers an increase of the yield by about 4–5 is possible. Very high SE yields ($\delta > 20$) are found on low density layers of KCl, BaF$_2$, (Goetze et al. 1964; Goetze 1968; Seiler and Stark 1965a).

In the SEM the energies of the PE are normally higher than $E_{PE}^m$. From $E_{PE} = 10$ to 100 keV, $\delta$ decreases with $E_{PE}^{-0.8}$ (Reimer et al. 1968). For Al, $\delta$ decreases from $\delta \approx 0.3$ ($E_{PE} = 10$ keV) to $\delta \approx 0.03$ ($E_{PE} = 100$ keV). Furthermore the SE yield in a SEM with a normal high vacuum is influenced by the growth of a contamination layer at the impact point of the electron beam (Wittry 1966; Seiler 1967). Figure 7 shows the way in which the SE yield of an Al surface decreases with bombarding time.

D. Escape depth of SE

Only SE excited near the surface can reach and escape from the surface (Copeland 1940). Normally it is assumed that the escape probability for SE produced at a distance $x$ from the surface, decreases with $e^{-x/\lambda}$, where $\lambda$ is the mean escape depth. Some measurements give the maximum escape depth $T$, other the mean escape depth $\lambda$ (Seiler 1967). Layers of increasing thickness were produced on a bulk substrate [see Fig. 7 (Seiler and Stähr 1965b)]. Beyond the particular thickness $T$ of the layer, $\delta$ no longer depends on the underlaying bulk material. If $\eta_{bulk} = \eta_{layer}$, $T$ gives a maximum escape depth of the SE and we can assume that $T \approx 5 \lambda$ (Seiler 1967). The escape depth of the SE from metals is of the order of $\lambda \approx 0.5$–1.5 nm, $T \approx 5$ nm; of insulators of $\lambda \approx 10$–20 nm, $T \approx 75$ nm. Mean escape depths of metallic oxides as Al$_2$O$_3$ and MgO and alkali halides as BaF$_2$, NaCl, and KCl are calculated by Kanaya et al. (1978). The high SE yield of insulators can be explained by the large escape depths of the SE. According to calculations by Ono and Kanaya (1979) the escape depth of SE can be correlated with regularities of the elements of the periodic table.

E. SE yield as a function of the angle of incidence

$\delta$ increases with increasing angle of incidence $\vartheta$ measured relative to the surface normal for $\vartheta < 80^\circ$ according to

$$\delta(\vartheta) = \delta_{\vartheta}(\cos \vartheta)^{-n} \quad \Delta \vartheta = \vartheta (\vartheta = 0').$$

(2)

The value $n = 1$ is valid for material with about $Z = 30$. For light elements $n$ increases (Bronshstein and Dolini 1968; Reimer and Pfefferkorn 1977) to about $n \approx 1.3$, and for heavy elements $n$ decreases to about $n \approx 0.8$. Kanaya and Kawakatsu (1972) give values 1.3 < $n$ < 1.5.

The increase of $\delta$ with increasing $\vartheta$ is due to the small escape depth of the SE. The longer the penetration distance of the PE within the escape depth of the SE, the higher the yield. With increasing $\vartheta$, $\delta''$, $E_{PE}^m$, and $E_{PE}^{\mu}$ increase (Bruining 1954; Kollath 1956; Reimer and Pfefferkorn 1977; Salehi and Flinn 1981). (See Fig. 8.) Using single crystals there is a fine structure superimposed on the monotonic increase of $\sigma$, $\eta$, and $\delta$ with increasing $\vartheta$ as shown in Fig. 9. This gives the

FIG. 8. Dependence of $\delta$ on the energy of the PE for different angle of incidence.

FIG. 9. Dependence of $\sigma = \delta + \eta$ on the angle of incidence for a Si (111) face (Kuhnle and Seiler 1970).
electron channeling pattern (ECP) in the SEM and the crystal orientation contrast in both in the SEM and the electron emission microscope using SE (Carle and Seiler 1966).

Crystallographic contrast effects in the SE, the RE, the Auger electron, and the x-ray yield can be explained by excitation of Bloch waves (Howie 1974). The fine structure of the SE and RE, i.e., the difference in the intensities $\Delta \delta$ and $\Delta \eta$, when changing the angle of incidence from a direction parallel to the lattice plane to the Bragg angle, was measured by a number of authors, e.g., Palmberg (1967), $\Delta \alpha/\alpha$, $\Delta \eta/\eta$, and $\Delta \delta/\delta$ decrease with increasing $E_{PE}$ as shown in Fig. 10 (Seiler and Kühnle 1970).

F. Contribution of backscattered electrons to SE yield

SE are not only released by PE but also by RE. This contribution by RE to the SE yield has been investigated by several authors [Everhart 1958; Kanter 1961; Bronshtein and Fraimont 1969; Seiler 1967, 1968a; Drescher et al. 1970; Kanaya and Kawakatsu 1972; Fitting et al. 1976; Reimer and Drescher 1977; Robinson 1974]. The SE yield is given by two parts:

$$\delta = \delta_{PE} + \eta \delta_{RE} = \delta_{PE}(1 + \beta \eta).$$  \hspace{1cm} (3)

$\delta_{PE}$ is the number of SE released per PE, $\delta_{RE}$ is the number of SE released per RE, $\beta = \delta_{RE}/\delta_{PE}$. For $E_{PE} > E_{RE}$ there is a higher probability for a SE to be excited by a RE than by a PE, so that $\beta > 1$. Bronshtein [reviewed by Seiler (1967)] measured $\beta \approx 4$ for $E_{PE} < 5$ keV, Drescher et al. (1970) measured $\beta \approx 2$ for $E_{PE} > 10$ keV. There are two reasons why $\beta > 1$. First, the mean energy of the RE is less than $E_{PE}$ and so $\delta$ is nearer to $\delta''$. Second, the lower average emergence angle of the RE is more favorable for the excitation of SE than the normal incidence of the PE beam.

This contribution of RE to $\delta$ is very important in the SEM. In such a case, where a contamination layer is on the object surface, image contrasts are caused by variation in $\eta$ rather than in $\delta$. For an Al surface with a contamination layer we get a yield $\delta \approx 0.10$ at $E_{PE} = 20$ keV. With $\beta = 2$ we can calculate that $\delta_{PE} = 0.07$ and $\delta_{RE} = 0.14$.

$\beta$ decreases with angle of incidence of the PE. Figure 11 shows the dependence of $\beta$ on the angle of incidence according to the measurements of Bronshtein and Denisov (1967) [surveyed by Seiler (1967)].

G. SE emission on the exit surface of thin transparent films

The SE are not only released on the surface of bulk material and thin foils but also on the back side (bottom) of transparent films if the energy of the PE is great enough (Fig. 12). There are many measurements on SE in transmission on thin films (Kanter 1961; Jahrreiss 1972; Hasselbach 1975; Reimer and Drescher 1977; Ono and Kanaya 1979; Kadlec and Eckertova 1970; Fitting 1974). On the entrance side the number $\eta \delta_{RE}$ of SE released by RE is given by

$$\eta \delta_{RE} = \delta_{PE} \beta \eta.$$  \hspace{1cm} (4)

On the exit side the number $t \delta_{E}$ of SE released by transmitted electrons can be expressed by (Reimer and Drescher 1977)

$$t \delta_{E} = \eta \delta_{RE}.$$  \hspace{1cm} (5)

FIG. 11. Decrease of $\beta$ with increasing angle of incidence according to the measurements of Bronstein (1965, 1967) for $E_{PE} = 4$ keV.

FIG. 12. SE emission of a transparent foil.
\[ t \delta_r = \delta_{PE} \beta_t \gamma, \quad \text{with} \quad \beta_t = \delta_t / \delta_{PE}. \quad (5) \]

\( t \) is the transmission coefficient, \( \delta_t \) is the number of SE/number of transmitted electron.

If the thickness of the foil is an appreciable fraction of the electron penetration depth, then the energy loss of the PE within the foil must be taken into consideration. At the exit side the maximum of the yield is at a higher value of \( E_{PE} \) than at the entrance side of the foil. Also the shape of the curve \( \delta_r \) \( (E_{PE}) \) is different from \( \delta(E_{PE}) \) because of the broadening of the energy distribution of the electrons in the foil. Figures 13 and 14 show the SE yield of Al foils on the entrance and on the exit surface for different energies of the PE as a function of the foil thickness. According to Reimer and Drescher (1977), \( \beta_t \) increases with \( D / R \), \( D = \) thickness of the foil and \( R = \) range of the PE. For \( D / R = 1 \) we get \( \beta_t = 3 \) for Al and \( \beta_t = 2 \) for Au. This can be explained similarly as in the case of RE. The transmitted electrons have lower energy than the PE and so their energy is nearer to the energy in the maximum of the SE yield and the angular distribution of the scattered transmitted electrons causes a higher SE yield than that of the PE at normal incidence. There is no experimental evidence that the conservation of momentum causes a higher SE yield in the direction of the PE beam as proposed by Robinson (1975). Within the solid an isotropic distribution of the excited electrons is assumed. Only in insulators with internal electric fields may there occur deviations from isotropic distribution.

**H. Noise in SE emission**

The incident PE beam is modulated by "shot noise." The noise in SE emission was investigated by several authors with respect to the SEM: Everhart (1958, 1970), Reimer and Pfefferkorn (1977), Baumann and Reimer (1981). According to Filipov (1966) for \( E_{PE} < 250 \text{eV} \) the SE emission shows a Poisson distribution. For energies used in the SEM the distribution of the SE yield is more complicated (Kurrelmeyer and Hayner 1937; Everhart et al. 1959; Reimer 1971) because the primary electrons and the RE can produce further SE at the surface when leaving the specimen. The distribution is therefore not simply a Poisson distribution but shows an increased relative variance. (Reimer and Pfefferkorn 1977).

**I. Exit area of excited SE**

The width of the exit area of the primary excited SE is determined by the mean escape depth \( \lambda \) of the SE. [See Fig. 15 (a)] (Wells 1957; Everhart 1958; Everhart and Chung 1972). If \( d_o \) is the diameter of the incident beam, the exit

![Graph](https://example.com/graph1.png)

**FIG. 13.** SE yield \( \delta \) at the entrance surface of a thin Al foil depending on the foil thickness for different PE energies according to Reimer and Drescher (1977).

![Graph](https://example.com/graph2.png)

**FIG. 14.** SE yield \( \delta \), at the exit surface of a thin Al foil depending on the foil thickness for different PE energies according to Reimer and Drescher (1977).

![Graph](https://example.com/graph3.png)

**FIG. 15.** Spatial distribution of SE released at a surface by PE and by RE. (a) The SE are released by the PE within a circle determined by the mean escape depth of \( \lambda \) of the SE. (b) Schematic intensity distribution of the SE (1) signal and SE (2) signal depending on the distance \( r \) from the impact of the primary beam. FWHM: Full width at half maximum.
diameter of the primary excited SE may be estimated by \(\sqrt{d_o^2 + \lambda^2}\). The SE excited by the RE will emerge over approximately the same area as the RE. The width of the exit area of these SE is determined by the range of the electrons [see Fig. 15(a)]. The schematic intensity of the both parts of the SE released by PE and by RE is shown in Fig. 15(b). The spatial distribution of SE released by a fine electron probe was measured by Hasselbach (1973), Hasselbach and Rieke (1978, 1982), with a combination of a SEM and an electron emission microscope (EEM). Normally in the EEM (Mollenstedt and Lenz 1963) the whole object surface is stimulated to emit electrons by UV light, heating, or by bombardment with PE or ions. It is possible to image the specimen with the emitted slow electrons by a cathode or immersion lens (see Fig. 16). This is a combination of an electric accelerating field and an electrostatic or magnetic electron lens. By using a cathode lens it is possible to get a picture of the spatial distribution of the SE emitted by a fine PE probe. The RE are prevented from reaching the image plane by an aperture in the focal plane of the cathode lens. According to Hasselbach and Rieke (1982) the spatial distribution of SE released by RE can both be approximated by Gaussian distributions. In Fig. 17 the full width at half maximum (FWHM) of the density distribution of SE released by RE on Si and Au is shown for energies of the PE from 20–70 keV.

**J. Influence of very thin surface layers on SE emission**

The work function \(\Phi\) has less effect on SE emission as compared with photo, thermionic- and field-electric emission. Metals with high \(\Phi\) often show also high \(\delta\). Of the three processes which contribute to SE emission, (1) electron production within the material, (2) migration of the electrons to the surface, and (3) escape of the SE over the surface potential barrier, the first two processes strongly depend on the bulk properties, while the third process should cause an increase in \(\delta\) with decreasing \(\Phi\). Palmberg (1967), Schäfer and Hölzl (1972) tried to reduce \(\Phi\) by evaporating Na on Ge or Pt surfaces. Reduction of \(\Phi\) from 4.79 to 2.3 eV increased \(\delta^m\) from 1.2 to 3.6, while \(E_{PE}\) increased from 700 to 2000 eV.

The SE emission can be influenced by resonant tunneling (Grundner and Halbritter 1980; Halbritter 1982, 1983). Thin absorbed layers of \(\text{H}_2\text{O}\) may enhance the SE yield up to 50% whereas the growth of unsaturated polymerized hydrocarbon layer may reduce the SE yield.

**K. Semiempirical theory of SE emission**

The elementary theory developed by Salow (1940) and Bruining (1954) has been reviewed by Dekker (1958) and Kollath (1956). The SE yield can be written in the following form:

\[
\delta = \int n(x, E_{PE}) f(x) dx.
\]

\(n(x, E_{PE})\) is the number of SE produced at a distance \(x\) from the surface by a PE of the energy \(E_{PE}\). \(f(x)\) is the probability that a SE produced at \(x\) reaches the surface and is emitted into vacuum.

Assuming that \(n(x, E_{PE})\) is proportional to the average loss per unit path length

\[
n(x, E_{PE}) = -\frac{1}{\epsilon} \frac{dE}{dx}.
\]

\(\epsilon\) is the energy required to produce a SE. It is generally assumed that

\[
f(x) = B e^{-x/\lambda},
\]

where \(B\) is a constant <1 and takes into account that only a fraction of the excited electrons migrates toward the surface and the probability of these electrons reaching the surface and passing over the surface barrier into vacuum.

According to Young (1956, 1957) the energy dissipation of PE within the material is approximately constant and hence

\[
\frac{dE}{dx} = \frac{E_{PE}}{R},
\]

where \(R\) is the range of the PE. From Eqs. (6)–(9) we get an expression for \(\delta (E_{PE})\)

\[
\delta = \int_0^{\infty} \frac{B E_{PE}}{\epsilon R} e^{-x/\lambda} dx,
\]
\[ \delta = B \frac{E_{PE} \lambda}{\epsilon} \left[ 1 - e^{-R/\lambda} \right]. \]  

(11)

Using the energy-range relation (Young 1959)

\[ R = \frac{1.15 \times 10^5 \left( \frac{E_{PE}}{p g/m} \right)^{1.35}}{\rho kg/m^3} [keV], \]

(12)

we get \( \delta = \delta(E_{PE}) \) or \( \delta = \delta(R) \). \( \delta(E_{PE}) \) is a function which increases with \( E_{PE} \) up to a maximum \( \delta^m \) at \( E_{PE}^m \) and then decreases.

The reduced yield \( \delta/\delta^m \) as a function of \( E_{PE}/E_{PE}^m \) is independent of the constants \( B, \epsilon, \rho \), which are characteristic for the material under consideration. Thus according to this theory the reduced yield curves should all follow a single universal curve.

\[ \frac{\delta}{\delta^m} = 1.11 \left( E_{PE}/E_{PE}^m \right)^{0.35} \left[ 1 - e^{-2.3 E_{PE}^{1.35}} \right], \]

(13)

with \( E_{PE} = E_{PE}/E_{PE}^m \).

Calculating the maximum of the yield curve \( \delta = \delta(R) \) we get the maximum for

\[ R = 2.3 \lambda. \]

(14)

According to several authors (Seiler 1967; Simon and Williams 1968; Buchholz 1969) the maximum of the yield curve is reached if the range \( R \) of the PE is approximately equal to the escape depth of the SE. For values \( R \) between maximum and mean escape depth, the theoretical value corresponds well with the experimental data. For \( E_{PE} \gg E_{PE}^m \) we get from Eq. (13)

\[ \delta = 1.11 \frac{\delta^m}{E_{PE}/keV} \left[ 1 - e^{-2.3 E_{PE}^{1.35}} \right]. \]

(15)

Using the experimental values of Ono and Kanaya (1979)

\[ \frac{\delta^m}{E_{PE}/keV} \approx k, \quad 1.87 \leq k \leq 2.4, \]

we get \( k = 2.1 \)

\[ \delta = 0.86 \left( \frac{E_{PE}}{keV} \right)^{0.35} \left[ 1 - e^{-2.3 E_{PE}^{1.35}} \right]. \]

(16)

Calculated values of \( \delta \) for \( E_{PE} = 20 \text{ keV} \) are shown in Table I. Those values are greater than experimental values, but are useful for estimating image contrasts. Seiler (1982) surveyed modern theories of SE emission.

II. SE IN THE SCANNING ELECTRON MICROSCOPE

In a SEM a fine PE beam is scanned over the specimen. The electron current leaving the surface is amplified and is used to modulate the brightness of a cathode ray tube, whose electron beam is moving over the screen synchronously with the PE beam. Surfaces were imaged using SE in experimental SEM's by Knoll (1935). Knoll and Theile (1939) show magnified images of surfaces with material, topography, and crystallographic contrast. Ardenne (1940) described a SEM for surface imaging. A voltage contrast was observed by Knoll (1941). Smith et al. (1955) used a SEM with a resolution better than 50 nm. Oatley and Everhart (1957) examined \( p-n \) junctions with a SEM.

A. SE detectors in the SEM

In the SEM the generated SE must be collected and the signal must be amplified without introducing additional noise.

1. Measurement of the sample current

The sample current \( i_s = i_{PE} - i_{SE} - i_{RE} \) can be measured by amplifying the voltage drop on a resistor or with a low noise operational amplifier with high input impedance (Knoll 1935; Oatley 1981). This signal gives, however, a poor \( S/N \) because for \( \sigma = 1, i_s = 0 \) and for \( \sigma > 1 \) the wanted signal \( i_{SE} \approx i_{PE} \).

2. Measurement of the emitted electrons with a collector

With a positive collector with a large solid angle of collection the collector current \( i_c = i_{RE} + i_{SE} \) can be measured. With a small solid angle of collection only those RE are measured which can be deflected by the positive potential toward the collector. With a small negative potential at the collector we get \( i_c = i_{RE} \).

3. Open multiplier and channeltron

SE were collected using an electron multiplier with Cu-Be dynodes by Smith and Oatley (1955). The use of a channeltron was discussed by Hantsche and Schreiber (1970). Problems are especially the change of the yield with time and the limited lifetime.

4. Everhart–Thornley detector (Fig. 18(a))

For SE images in the SEM mostly a scintillator/photo-multiplier is used (Everhart 1958; Everhart and Thornley 1960). It consists of a scintillator at the end of a light pipe, surrounded by a shielding cage with a metal grid at one end. The scintillator is covered with a very thin layer of aluminum and held at about \( +10 \) kV. The cage is directed towards the specimen and held at some potential: with a potential of \( +200 \) V most of the released SE in the SEM are collected. With a potential of \( -100 \) V only the RE which are emitted in the direction of the collector are measured. The light pipe ends at a vacuum window, pointing to a photomultiplier tube. To collect the SE the specimen is tilted towards the cage and the SE are drawn to the cage and towards the collector. Most of them travel through the grid, are accelerated to the coated scintillator, pass through the aluminum film, and create photons. This light is conveyed by the light pipe to the photomultiplier and converted into electrical signals.

According to Everhart (1958, 1959), 65% of the signal stems from SE, 5% from RE, and 30% from SE released by RE at the walls of the SEM. This detector is suitable for currents in the range of \( 10^{-16} - 10^{-15} \) A. Light pipe and vacuum feedthrough losses typically reduce the signal to 35% of the created photons.

The noise of different detection systems for SE and RE using a scintillator/photomultiplier combination was studied by Baumann and Reimer (1981). Very interesting are the investigations by Austra et al. (1978, 1982) to use single
crystal yttrium aluminum garnet (YAG) activated with cerium as scintillator for SE and PE. The advantage is its high relative efficiency; this involves losses due to optical matching, spectral transmittance of the light pipe, losses on the photocathode of the multiplier, and losses due to insufficient technological optimization of the scintillator.

The SE emitted from the specimen travel different trajectories in the electric field between the lens and the collector according to their different initial energies and momenta. Figure 19(a) shows the trajectories of SE leaving the object surface with energies from 0 to 10 eV towards the collector at +200 V. Figure 19(b) shows the deflection of SE starting of an object at different potentials from −15 to +15 V (Everhart 1958, 1968). Computation of trajectories in voltage contrast detectors were published by Kursheed and Dinis (1983).

5. Detector for imaging the surface only with SE released from a small sample area [Fig. 18(b)]

This detector by Hasselbach et al. (1983) is similar to the arrangement for measuring the radial spread of the SE (Sec. I 1). The cathode lens gives an enlarged image of the specimen in the plane of the fluorescent screen. The aperture “a” in the focal plane of the cathode lens prevents the RE from reaching the image plane. The aperture “b” selects the circle plane of the object surface contributing to the signal at the scintillator. The deflection element is driven in synchronism with the scanning of the PE beam in order to compensate for the deflection of the PE beam. OnlySE released in a small area around the PE beam are focussed on the scintillator, so contrast may be increased.

6. Detector for quantitative measurement of voltage contrast [Fig. 18(c)]

The voltage contrast can be measured in the SEM by passing the SE through an electrostatic energy analyzer between specimen and the electron detector (Wells and Bremer 1968). Another arrangement consists (Menzel and Kubalek 1981) of an immersion or cathode lens for the generation of a high extraction field, a 127° analyzer, a retarding
field analyzer and an Everhart–Thornley detector. It allows the measuring of small potential differences between adjacent object details (see Voltage II contrast).

B. SE signal in the SEM

In the conventional SE imaging mode in the SEM several different SE signals reach the detector: see Fig. 2 (Schar et al. 1967; Seiler 1968a; Reimer and Pfefferkorn 1977; Peters 1982a,b).

SE (1) signal: SE produced only by PE: \( \delta_{\text{PE}} \);
SE (2) signal: SE produced by RE at the surface of the specimen: \( \eta \delta_{\text{RE}} \);
SE (3) signal: SE produced by RE at the walls of the SEM: \( \alpha \eta \delta_{\text{walls}} \);
RE (4) signal: RE which reach the detector because they are emitted in this direction.

The SE (1) signal is the desired signal for high resolution, because the SE are emitted at the impact point of the PE beam. This signal is however very sensitive to the conditions of the surface. In a normal SEM we often get a contamination layer at the surface and so the material contrast is poor, compared with topography contrast. In general the SE (2) signal may decrease the lateral resolution because of the lateral spread of the RE. This signal gives however high material contrast because often differences in \( \eta \) are greater than in \( \delta \). Moreover, this material contrast is nearly independent of thin contamination layers. This signal also makes it possible to see details within the specimen far beyond the escape depth of the SE. This will be discussed in Sec. II H (Information Depth).

The SE (3) signal may increase the SE (2) signal. This is useful to get a high material contrast. Reimer and Volbert (1979) used a negatively biased MgO coated converter plate below the pole piece of the final lens to increase this SE (2) signal.

The SE (3) signal decreases the lateral resolution like the SE (2) signal. In order to decrease the SE (3) signal a RE-absorption plate (a carbon coated aluminum plate with a very low SE yield) can be used (Peters 1982a,b) or a positively biased converter plate below the pole piece of the final lens to prevent SE released by the RE from leaving this plate (Bau mann and Reimer 1981). The RE (4) signal depends on the position of the collector relative to the object. It can be prevented using special detectors with a cathode lens (immersion objective) (Hasselbach and Rieke 1982; Menzel and Kubalek 1981). In the cathode lens the slow SE are focussed by an electric field. An aperture in the focal plane of the cathode lens prevents the RE from reaching the collector.

A further SE signal was shown by Hasselbach and Rieke (1976) on objects with sharp edges using a SEM with a special detector with a cathode lens. If the PE beam hits the specimen close to sharp edges, the scattered primaries leaving the edges may again hit the specimen far from their original impact point contributing again to the SE yield (Wells 1978).

The normal SE signal in the SEM is a mixture of different contrast mechanisms. The use of multidepctor system makes it possible to isolate one wanted signal. For imaging the specimen surface the SE (1) signal is wanted, the SE (2) signal confuses the detection of surface properties. The SE (2) signal is however proportional to \( \eta \). So if we measure the RE signal \( \eta \) with another detector we can perform a subtraction: SE signal minus \( k \eta \), so we might be able to select a value of the constant \( k \) to enhance surface details by producing a signal which approximately consists only of SE (1) signal (Crewe and Lin 1976; Reimer 1982; Volbert 1982).

C. Changes of SE emission with time

In the SEM, the SE yield may change with time. In normal vacuum systems a contamination layer can be built by PE and also by RE. The formation of insulating films under electron or ion bombardment was discovered by Stewart (1934). Several papers have been written on this subject and have been summarized by Hren et al. (1979) and Love et al. (1981). Figure 7 shows the decrease of the SE yield of an Al target depending on the thickness of the contamination layer. With fine electron probes contamination rates of 50 nm/s are possible.

In the SEM the contamination is to be seen as a dark area when the magnification is decreased. On thin foils the contamination may increase the brightness of the field under investigation as shown by Martin (1971).

Changes in SE yields, depending on electron beam dose may also occur under UHV conditions (Le Gressus et al. 1979).

The electron beam can produce electron stimulated desorption and alter the surface composition of the investigated object. In insulators, as for example in glasses, the electron beam breaks bonds and forms ion-electron pairs and raises the temperature of the material. Both effects cause an increase in alkali ion mobility. During the electron bombardment an electron field within the insulating objects builds up and forms a diffuse negative charge layer at a depth up to the range of the PE. This electric field can cause a field assisted alkali ion diffusion (Ohuchi et al. 1980). Specimen charging is a serious problem in the SEM. A summary of charging effects relating to the internal electrical fields and especially to the external fields was given by Shaffner and Hearer (1976).

D. Contrast in the SEM

Different SE yields of adjacent object elements cause different brightness between the corresponding picture elements (Knoll and Theile 1939; Smith and Oatley 1955; Wells 1957; Everhart et al. 1959). The difference in intensity \( \Delta I \) of object elements \( A \) and \( B \) must be greater than the signal/noise \( (S/N) \). In this part only the contrast of large parts of the specimen will be estimated. The contrast is given by

\[
C = \frac{I_A - I_B}{I_A + I_B} = \frac{\Delta I}{2I},
\]

with

\[
T = (I_A + I_B)/2.
\]

There are different types of contrast in the SEM (Only contrasts due to differences in \( \delta \) are considered):
(1) Material contrast due to different materials at the surface.

(2) Topography contrast due to different inclinations of surface details.

(3) Electron channeling and crystallographic contrast due to different crystallographic orientation with respect to the PE beam.

(4) Voltage type I contrast due to different δ for different energies of the PE at electrically floating object details.

(5) Voltage type II contrast due to different potentials at surface details often combined with (6).

(6) Voltage type III contrast due to the deflection of SE in the electric fields at the surface.

(7) Voltage contrast at the surface of ferroelectrics.

(8) Magnetic type I contrast due to the deflection of SE in magnetic fields at the surface.

1. Material contrast

   a. Clean surfaces. This contrast caused by surface materials with different SE yields. The difference in intensity ΔI is at its maximum for material A with δ A 0 E A 0 and material B with δ B 0 E B 0 when the energy of the PE is either E A 0 or E B 0 . These energies are normally not used in SEM because of the low brightness of the electron guns at low PE energies and the high contamination rates. According to formula (16) the SE yield can be estimated for 20 keV.

   \[
   C = \frac{\left( \delta_{A}^{1.35} - \delta_{B}^{1.35} \right)}{\left( \delta_{A}^{1.35} + \delta_{B}^{1.35} \right)}. \tag{18}
   \]

   For an object consisting of Ti: δ A = 0.8 and Pt: δ B = 1.35 we get with SE E PE = 20 keV: C1 = 0.34; and with RE Ti: η = 0.3, Pt: η = 0.43 we get C 2 = 0.18.

   b. Contaminated surfaces. In this case we must consider in Eq. (3) if the difference between δ A and δ B is caused by different δ PE or by different η. In the first case η A = η B ; δ A = δ B PE the contrast disappears quickly with increasing thickness of the contamination layer C 1 = 0.

   In the second case η A ≠ η B ; δ A = δ B PE we have a material contrast even with a contamination layer.

   \[
   C_2 = \frac{\beta (\eta_A - \eta_B)}{2 + \beta (\eta_A + \eta_B)}. \tag{19}
   \]

   For Ti: η = 0.3 and Pt: η = 0.43 we get with β = 2: C 2 = 0.08. In this case, which we have normally in the SEM, the contrast is less than using RE. However in a UHV-SEM we can get a higher contrast using SE, especially with low energy PE.

2. Topography contrast

The topography contrast is caused by the dependence of δ on the angle of incidence [Fig. 20(a)]. Object details with different inclinations δ and (δ + Δδ ) with respect to the PE beam are imaged with different brightness. From Eq. (2), δ = δ 0 (cos δ ) − n follows for n = 1 for the contrast between the planes with the inclination Δδ

\[
C = \frac{\cos \theta - \cos(\theta + \Delta \theta)}{\cos \theta + \cos(\theta + \Delta \theta)}/\left[ \cos \theta + \cos(\theta + \Delta \theta) \right] \tag{20a}
\]

for θ ≠ 0 follows

\[
C_1 = \frac{1 - \cos \Delta \theta}{\cos \theta + \cos(\theta + \Delta \theta)} \tag{20b}
\]

for two planes with a fixed small inclination Δδ we get for θ ≠ 0

\[
C_2 = \frac{\Delta \theta}{2} \tan \theta. \tag{20c}
\]

So a small topography contrast due to small differences Δδ can be enhanced by tilting the object to higher values of δ (Everhart 1958): for δ = 0°, Δδ = +10° results in C = 0.007; for δ = 50°, Δδ = +10° results in C = 0.125. This is valid for a surface with only small angular variations. If the surface is rough, the collection efficiency of the collector must be considered too.

In deep notches the field strength of the electric field...
between specimen and collector is sometimes too small to collect all emitted SE. The electrical field strength at a smooth surface is less than 100 V/cm if a collector potential of 250 V is used. In deep notches the electrical field is smaller and so the image contrast depends on both the energy distribution and the angular distribution of the emitted SE [Fig. 20(b)]. This shadowing effect is however not so strong as with RE.

At edges we get a very high SE yield [Fig. 20(c)]. One contribution is given because the path length of the PE within the escape depth of the SE along the edge is large. More important is the effect that the edge can be within the scattering cloud of the PE and RE. This results in a high SE yield. This part of the edge effect depends on the range of the PE. Fine object structures at the surface are better resolved at low PE energies.

### 3. Voltage contrast

**Voltage type I contrast** [Fig. 21(a)]. The SE yield depends on the energy of the PE striking the surface. Different energies of the PE at the surface may not arise only due to different beam energies, but also to different potentials of object details. This can be caused by charging of electrical floating object details. As shown in Sec. 1 C the second crossover $E_{EPE}$ in the curve $\delta$ depending on $E_{PE}$ (see Fig. 6) represents a stable equilibrium. The different potentials of object details can be produced with an additional electron beam. Image contrast in SEM can be used for a contactless testing of electrical networks as shown by Pfeiffer et al. (1981), Brünger (1982), Pfeiffer (1982), and Hohn et al. (1982).

**Voltage type II contrast** [Fig. 21(b)]. The energy distribution of SE is fixed relative to the potential of the surface. A change of the surface potential causes a shift of the energy spectrum of the SE. Recording the energy distribution of the emitted SE with an energy analyzer gives a possibility to measure the surface potential (Everhart 1958; Wells and Bremer 1968; Menzel and Kubalek 1981). The basis of quantitative voltage measurements is the fact that all electrons of the SE spectrum emitted from an electrode at a certain potential gain the same kinetic energy towards the detector. In this way the whole spectrum is linearly shifted with the applied specimen voltage. The linear shift of the maximum of the spectrum is then a direct measure of the applied specimen voltage. According to Gopinath (1977) the voltage resolution, i.e., the minimum measurable voltage in the SEM, for retarding field analyzers depends on the signal-to-noise ratio, the bandwidth of the detection system, and the primary electron beam current. The detection of voltage differences of about 1 mV is possible (Menzel and Kubalek 1979).

**Voltage type III contrast.** This may be caused by the deflection of the slow SE by electric fields parallel to the surface between different object details. This is similar to the magnetic type I contrast in SEM.

**Voltage type IV contrast.** The domain structure of ferroelectrics gives a contrast in SEM. This may partially be due to voltage III contrast. On the basis of a surface layer, Uchikawa (1982) presented a theory which incorporates the effect of local electrical fields produced across the surface layer by the underlying spontaneous polarized charge onto the SE yield.

### 4. Magnetic type I contrast

A contrast in SEM pictures may arise from the influence of magnetic fields of the specimen. The magnetic fields especially those parallel to the surface can deflect the slow SE so that they cannot reach the detector (Gentsch and Reimer 1972). The magnetic type II contrast shows the influence of internal magnetic fields, deflecting the PE and RE within the specimen.

### 5. Electron channeling and crystallographic contrast

The monotonic increase of $\delta$ with increasing angle of incidence away from the surface normal is only to be seen on amorphous or microcrystalline surfaces. On single crystal targets there are maxima and minima superimposed to the monotonic increase (Fig. 9). This causes the ECP in SEM (Joy et al. 1982). ECP can be obtained in the SEM with either SE or RE. The contrast of ECP, registered with SE, depends very sensitively on thin amorphous layers or contamination at the surface. The Bloch waves (Sec. I E) are strongly attenuated with increasing depth, and the ECP are formed in a layer near the surface. So the energy of those RE which give the ECP is near to the energy of the PE and so $\beta$ is smaller than usual (Seiler et al. 1970; Seiler 1976). This is in accordance with the high contrast of the ECP in “Low Loss” SEM (Wells 1971).
E. Information depth in the SEM using SE

The information depth \( D \) is defined as the depth below the surface of the object contributing to the SEM picture. Object details in a depth \( d \), with \( d < D \), can be recognized. \( D \) depends on the minimal contrast which is detectable in the SEM, on the maximum exit depth of RE, even if SE are used for imaging, and on the difference \( \Delta \eta \) in the backscattering coefficients between the object detail and the adjacent material (Seiler 1976; Wells et al. 1982). Figure 22 shows an object detail of material \( B (\eta_B) \) embedded in material \( A (\delta, \eta_A) \) in a depth \( d \).

The intensity in the object detail \( A \) is

\[
I_A = I_{PE} \delta_{PE} + I_{PE} \eta_A \delta_{RE}.
\]

(21)

The intensity at the surface of the specimen above object detail \( B \) is the sum of

\[
I_B = I_{PE} \delta_{PE},
\]

\( I_B \) = those SE, released by the RE backscattered in material \( A \),

\( I_B \) = those SE, released by the RE backscattered in material \( B \). With increasing thickness \( d \) of the layer the intensity of the PE decreases.

For a rough estimate of the information depth we assume that the number of penetrating electrons in a layer of the thickness \( d \) decreases with \( I_{PE} e^{-4d/R} \) with \( R \) the range of the PE. The maximum of the escape depth of the RE is about \( R/2 \) and the number of the RE according to their lower mean energy decreases in a layer of a thickness \( d \) with \( I_{RE} e^{-8d/R} \) (Seiler 1976). The difference \( \Delta I \) in intensities of the SE is

\[
I = I_{PE} \delta_{RE} (\eta_A - \eta_B) e^{-8d/R}.
\]

(22)

The contrast in the SEM for SE with \( \beta = 2 \) and the mean value \( \bar{\eta} = 0.25 \) is

\[
C = \frac{\Delta I}{2I_{PE}(\delta_{PE} + \bar{\eta} \delta_{RE})} \approx 0.67 (\eta_A - \eta_B) e^{-8d/R}
\]

(23)

and with Eq. (12) we get the information depth

\[
D = R/8 \ln [67(\eta_A - \eta_B)].
\]

(24)

FIG. 23. Information depth in the SEM as a function of \( E_{PE} \).

and with Eq. (12) we get the information depth

\[
\frac{D}{\mu g/cm^2} = 1.4 \ln [167 \Delta \eta] \left( \frac{E_{PE}}{keV} \right)^{1.35}
\]

(25)

Figure 23 shows these functions for various \( \Delta \eta \) which agree rather well with experimental results. Figure 24 shows an example for the dependence of the information depth on the energy on the PE. The estimate of the information depth is similar to the measuring of the thickness of layers of RE (Hohn et al. 1976; Wells et al. 1982). Calculations on the dependence of SEM contrast upon electron penetration were published by George and Robinson (1976).

F. Lateral resolution in the SEM using SE

(a) Signal/Noise ratio. In order to discriminate between adjacent picture elements by their different brightness, the difference in the signal quanta caused by the released SE must be high enough compared with the statistical fluctuations. A beam of \( 10^{-11} \) A is satisfactory for most objects (Oakley 1981; Broers 1982).

(b) The diameter of the PE beam. The beam diameter in which we can focus a beam depends on the primary beam energy, the brightness of the electron gun, the energy spread of the electron beam, the chromatic aberration coefficient of the final lens \( C_e \), the spherical aberration coefficient of the final lens \( C_s \), and the focal length of the final lens (Nagantani and Ohura 1977).

Figure 25 shows the minimal beam diameter for a beam with \( 10^{-11} \) A at 25 keV with \( C_e = 18 \) mm, \( C_s = 11 \) mm depending on the brightness of the electron gun with an energy spread of 0.3 eV typical of a field emission gun. Using thermal cathodes the energy spread is normally greater (Broers 1982).

(c) The spatial emission distribution of SE. The spatial emission distribution of the SE (1) signal is much smaller than the spatial emission distribution of the SE (2) signal and of the RE. If the SE (1) signal is strong enough compared with SE (2) signal the resolution of a SEM using SE is deter-
G. Conditions for high resolution in the SEM

To get a beam diameter of about 1 nm with a beam current of $10^{-11}$ A we need a field emission gun. High resolution images are possible with the SE (1) signal (Peters 1982a,b). The SE (3) signal can be suppressed by using an electron absorption device attached to the pole piece of the final lens or a positive converter plate. The main problem is the reduction of the SE (2) signal which has the spatial emission distribution of the RE which decreases the resolution. The use of a multi detector system works at the cost of the $S/N$. Especially for imaging of biological objects with low atomic number the SE (2) signal is weak if the coating metal film to prevent charging is as thin as possible and homogeneous (Evans and Ranks 1981). According to Peters (1982a,b) it is possible to get fine crystalline homogeneous films with a thickness of about 2 nm with the metals tantalum, niobium, and chromium. For these objects the quality of the images in the SE (1) image mode with a SEM with a field emission gun equals according to Peters (1982a) that of images obtained in the transmission electron microscope for identically coated specimens.

ACKNOWLEDGMENTS

The author thanks Dipl. Phys. H. E. Bauer and Dr. O. C. Wells for many helpful discussions, Professor Dr. T. E. Everhart for valuable hints, Mrs. Jacoby for very careful drawing of the figures, and Mrs. Paukert for patient typing assistance.

LIST OF ABBREVIATIONS

SE: Secondary electron
SEM: Scanning electron microscope
PE: Primary electron
SEE: SE emission
$\lambda$: Mean escape depth of SE
$T$: Maximum escape depth of SE
UHV: Ultrahigh vacuum
RE: Backscattered or reflected electrons
$E_{se}$: Most probable energy of the SE
HW: Full width at half maximum
$\delta$: SE yield
$\eta$: RE coefficient


