
**Surface chemical analysis —
Determination of surface elemental
contamination on silicon wafers by
total-reflection X-ray fluorescence
(TXRF) spectroscopy**

*Analyse chimique des surfaces — Détermination de la contamination
en éléments à la surface des tranches de silicium par spectroscopie de
fluorescence X à réflexion totale*



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Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

The procedures used to develop this document and those intended for its further maintenance are described in the ISO/IEC Directives, Part 1. In particular the different approval criteria needed for the different types of ISO documents should be noted. This document was drafted in accordance with the editorial rules of the ISO/IEC Directives, Part 2 (see www.iso.org/directives).

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For an explanation on the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO's adherence to the WTO principles in the Technical Barriers to Trade (TBT) see the following URL: Foreword - Supplementary information

The committee responsible for this document is ISO/TC 201, *Surface chemical analysis*.

This second edition cancels and replaces the first edition (ISO 14706:2000), which has been technically revised.

Introduction

This International Standard was prepared for the measurement of surface elemental contamination on silicon wafers on the basis of three existing standards: ASTM F 1526, SEMI Standard M33, and a UCS (Ultra-Clean Society) standard published by the Institute of Basic Semiconductor Technology Development.

TXRF needs reference materials to perform quantitative analyses. Certified reference materials are not available at low densities of 10^{10} atoms/cm². Even if they were available, the possibility of contamination from the environment reduces the shelf life of such reference materials.

Therefore, the TXRF reference materials are to be prepared and analysed for calibration by each relevant analytical laboratory. Thus, two standards, one for the TXRF measurement procedure and the other for the preparation of reference materials, are necessary. This International Standard concerns the former part.

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Surface chemical analysis — Determination of surface elemental contamination on silicon wafers by total-reflection X-ray fluorescence (TXRF) spectroscopy

1 Scope

This International Standard specifies a TXRF method for the measurement of the atomic surface density of elemental contamination on chemomechanically polished or epitaxial silicon wafer surfaces.

The method is applicable to the following:

- elements of atomic number from 16 (S) to 92 (U);
- contamination elements with atomic surface densities from 1×10^{10} atoms/cm² to 1×10^{14} atoms/cm²;
- contamination elements with atomic surface densities from 5×10^8 atoms/cm² to 5×10^{12} atoms/cm² using a VPD (vapour-phase decomposition) specimen preparation method (see 3.4).

2 Normative reference

The following documents, in whole or in part, are normatively referenced in this document and are indispensable for its application. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

ISO 14644-1, *Cleanrooms and associated controlled environments — Part 1: Classification of air cleanliness*

3 Terms and definitions

For the purposes of this document, the following terms and definitions apply.

3.1

total reflection

complete reflection of glancing X-ray radiation into a medium having larger X-ray optical density value at a boundary between two media

Note 1 to entry: The refractive index of silicon for X-rays is less than 1. X-rays which are incident on the surface at a small glancing angle are therefore, totally reflected from the surface at an angle equal to the glancing angle.

3.2

glancing angle

angle between the specimen surface plane and the virtual plane containing incident X-rays which impinge on the sample surface

3.3

critical angle

glancing angle corresponding to the first point of inflection in the plot of the sample matrix X-ray fluorescence against the glancing angle

3.4

VPD-TXRF method

method in which impurities on the surface are collected by the so-called VPD procedure, i.e. the non-volatile products formed by acid decomposition of the oxide at the wafer surface are collected by a droplet of collecting agent, usually ultra-pure hydrofluoric acid, and dried in a manner which gives the least environmental contamination, the residue from the droplet, subsequently being analysed by TXRF

3.5

spurious peaks

detected peaks that do not originate from impurities on the silicon wafer

Note 1 to entry: Spurious peaks are due to fluorescent X-rays originating from elements in the detector or the X-ray path. The fluorescent X-rays are excited by direct scattering or reflection of incident X-rays. This phenomenon leads to an increase in the measurement error. Spurious peaks seriously affect analytical measurements in the contamination range from ca. 10^{10} atoms/cm² to ca. 10^{11} atoms/cm².

4 Abbreviated terms

FWHM full width at half maximum

RM reference material

SSD solid-state detector

SDD silicon drift detector

TXRF total-reflection X-ray fluorescence

VPD vapour-phase decomposition

5 Principle

When a specimen is irradiated with X-rays, fluorescence X-rays at characteristic energies of the elements that constitute the specimen are generated. The intensities of the fluorescence X-rays are proportional to the amounts of each element in the specimen.

Total reflection of the incident X-rays on the specimen reduces a penetration depth of the incident X-rays in the specimen. This allows more selective excitation of the fluorescence X-rays from the surface region, including atoms deposited on the surface of the silicon wafer. Consequently, a spectrum of fluorescence X-rays with a large ratio of signal to background (S/B) and signal to noise (S/N) can be obtained.

The detection limit depends upon the atomic number, the excitation energy, the photon flux, the detector resolution and energy-dependent detection efficiency, the energy bandwidth of the excitation X-rays, the specimen-related shape and statistics of TXRF spectral background, the instrument-related noise magnitude, the integration time, and the accuracy of calibration of the RM and the blank value. For constant instrumental parameters, the interference-free detection limits vary over two orders of magnitude as a function of the atomic number of the analyte element.

NOTE The depth of measurement can vary with the glancing angle, but in the case of film-type contamination it is usually less than 5 nm. The area of measurement consists of a circle of ca. 10 mm in diameter, though it varies depending on the relative position of the X-ray detector and the specimen. In the case of particulate-type contamination on a clean surface, the yield of fluorescence X-rays varies depending on the sizes, distribution, and constituent elements of the particles.

6 Apparatus

6.1 The apparatus for TXRF shall include at least the following components: an X-ray source; a monochromator; a specimen stage capable of movement in three orthogonal directions; an X-ray detector (SSD or SDD); and a computerized signal-processing system.

6.2 X-rays which have been monochromatized shall be used as the incident X-rays.

6.3 The fluorescence X-ray detector shall have sufficient energy resolution to analyse the Mn-K-L_{II,III} line with an FWHM of 200 eV or less.

6.4 The specimen stage, which sets the glancing angle, shall be controlled to a reproducibility within $\pm 0,17$ mrad ($0,01^\circ$) in the range between 0 mrad (0°) and 8,7 mrad ($0,5^\circ$).

6.5 The atmosphere in the specimen chamber shall be able to be brought to a reduced pressure or replaced with helium gas or nitrogen gas, as required.

7 Environment for specimen preparation and measurement

7.1 The local environment (i.e. airborne particles, temperature, humidity) for specimen preparation and measurement shall be equal to or better than ISO Class 4 in accordance with ISO 14644-1.

NOTE The unwanted deposition of airborne particles which are composed of the elements that are being measured will cause an increase in the error of measurement.

7.2 The mechanical vibration at the location where the apparatus is installed shall be as small as possible and shall not be greater than 5×10^{-3} m/s² (0,5 Gal).

NOTE The mechanical vibration will degrade the energy resolution of the detection system, which will, in turn, degrade the detection limits and peak deconvolution.

8 Calibration reference materials

8.1 Calibration reference materials (RMs) used to establish a reliable calibration procedure shall consist of an RM on which known amounts of impurities have been deposited and a blank RM used to determine the level of contamination of the calibration RM (see [Annex A](#)).

8.2 The RMs shall be prepared from a chemomechanically polished wafer with a certain quantity of Ni or Fe uniformly deposited on its surface as the RM element. The atomic surface density of the RM element shall be ca. 1×10^{12} atoms/cm² to 1×10^{13} atoms/cm² (see [Annex C](#)).

8.3 How the RM element is located on the RM surface shall be verified by an anglescan (see [Annex E](#)).

8.4 The amount of RM element deposited on the surface of the wafer shall be determined by a reliable quantitative method of analysis.

8.5 The blank RM shall be a chemomechanically polished or epitaxial wafer. The magnitude of contamination of the surface region of the blank RM shall be below the detection limit for specified elements. The crystallographic orientation of the blank RM shall be the same as that of the RM.

8.6 The RM and the blank RM shall be stored in the same container.

9 Safety

This test method uses X-ray radiation. Consequently, it is absolutely essential to avoid exposing any part of the body to the X-rays produced by the apparatus. It is especially important to keep hands and fingers out of the path of the X-rays and to protect the eyes from scattered secondary X-ray radiation. Each country has its own safety regulations and requirements concerning X-rays. These shall be observed.

10 Measurement procedure

10.1 Preparation for measurement

10.1.1 All specimens shall be chemomechanically polished or epitaxial wafers.

10.1.2 For the VPD-TXRF method, the impurities on the surface are collected by VPD, i.e. acid decomposition in a droplet which is then dried in a manner that produces the least environmental contamination (see [Annex D](#)).

10.1.3 For a series of measurements, including the calibration measurements, the crystal orientation of the specimen on the specimen stage shall be the same for each specimen.

10.1.4 Set the glancing angle at between 25 % and 75 % of the critical angle.

NOTE 1 Due to the physical form of particulate-type contamination, an angle that is too low will cause a larger error.

NOTE 2 The critical angle is a function of the incident X-ray energy. It is 3,20 mrad (0,18°) for 9,67 keV (W-L_{II}-M_{IV}), 1,78 mrad (0,10°) for 17,44 keV (Mo-K-L_{II,III}) and 2,72 mrad (0,16°) for 11,4 keV (Au-L_{II}-M_{IV}).

10.1.5 Set the following parameters as specified.

- a) For a rotating anode, the excitation voltage of the X-ray source shall be 30 kV or more, with the excitation current set to 200 mA or more and the integration time set to 500 s or more.
- b) For a sealed Mo or W anode X-ray tube, the excitation voltage of the X-ray source shall be 30 kV or more, with the excitation current set to 40 mA or more and the integration time set to 500 s or more.
- c) If the X-ray intensity at the detector is too high for the detection system, the excitation current shall be adjusted to a lower value to give an appropriate count rate.

10.1.6 Move the centre of the specimen under the centre of the detector. While rotating the specimen, generate and measure fluorescence X-rays to find the azimuth with the least spurious peaks. When possible, the measurement hereafter shall be conducted at the same azimuth.

If the apparatus does not allow the best-fit azimuth to be set at off-centre positions so as to avoid spurious peaks, care shall be taken when evaluating the surface-mapping data.

10.2 Preparing a calibration curve

10.2.1 Measure the blank RM and determine the integrated intensity of the fluorescence X-rays generated by the blank RM.

10.2.2 Measure the RM under the same conditions as specified in [10.2.1](#). In the case of the VPD-TXRF method, place the residue from the RM under the centre of the detector.

10.2.3 Verify that the measured value of the integrated intensity of the X-rays generated by the blank RM is less than 10 % of the integrated intensity from the RM.

If the integrated intensity from the blank RM is 10 % or more of that from the RM, discard both the RM and the blank RM and prepare a new set comprising an RM and blank RM.

Carry out this check at appropriate intervals.

10.2.4 Determine the integrated intensity of the X-rays generated by the RM element by means of procedure a) and/or b) specified below.

- a) Smooth the measured numerical values by digital processing.

Determine the integrated intensity of the X-rays by subtracting the background from the measured numerical values.

- b) Determine the Gaussian function that best fits the measured numerical values. Then determine the integrated intensity from the peak height and the half-width of the Gaussian function.

10.2.5 Obtain a calibration curve (a plot of the atomic surface density versus the integrated X-ray intensity from the RM element). The curve shall pass through the origin.

10.3 Measurement of a test specimen

10.3.1 Measure the test specimen under the same conditions as specified in [10.1](#). If using the VPD-TXRF method, place the residue from the test specimen under the centre of the detector.

10.3.2 Determine the integrated intensity of the X-rays generated by contamination elements in the same manner as specified in [10.2](#).

When two or more fluorescence X-ray lines overlap, use the method of deconvolution to obtain the integrated intensity of the X-rays for the subject element.

NOTE 1 The repeatability and reproducibility of the measurement for the subject element will vary with the kind of X-ray used.

NOTE 2 Deviation of the glancing angle of the incident X-rays from the set value will increase the measurement error.

NOTE 3 Greater surface roughness will increase the measurement error.

NOTE 4 The values obtained from VPD residues will depend greatly on the physical form of the residue and the elements contained in the residue.

11 Expression of results

11.1 Method of calculation

By using Formulae (1) and (2), calculate the atomic surface density, C_m for each of the contamination elements from the results obtained in [Clause 10](#).

$$K = \frac{C_S}{I_S} \quad (1)$$

$$C_m = K \times \frac{I_m}{S_R} \quad (2)$$

where

K is the slope of the calibration curve obtained in [10.2.5](#);

C_S is the atomic surface density of the RM element, in atoms/cm²;

I_S is the integrated intensity of the fluorescence X-rays from the RM element, in counts per second (cps);

C_m is the atomic surface density of the contamination element on the test specimen, in atoms/cm²;

I_m is the integrated intensity of the fluorescence X-rays from the contamination element on the test specimen, in cps;

S_R is the relative sensitivity factor, which corrects for the difference in sensitivity for each element.

NOTE In order to determine values of the relative sensitivity factor, S_R for other elements, measurements are often made on two or more specimens with differing contents of these elements. The results can then be used to prepare calibration curves other than those obtained using theoretical relative sensitivity factors. The relative sensitivity factor can be determined in terms of these curves using the RM element or can be calculated as shown in [Annex B](#).

11.2 Blank correction

For measurements below 10¹¹ atoms/cm², where the instrumental blank is not negligible, the atomic surface density, C_m shall be corrected by subtracting the atomic surface density, C_0 of a fresh contamination-free specimen, with the same crystal orientation, measured under the same conditions as specified in [10.1](#) and [10.2](#) and calculated as described in [11.1](#).

12 Precision

An international inter-laboratory test programme based on the method specified in this International Standard was carried out by 15 laboratories from Japan, Europe, and USA. Four test specimens and one RM were distributed as one set. 17 sets of measurements were obtained from the 15 laboratories. Repeatability and reproducibility were calculated in accordance with the principles of ISO 5725-2.^[1] A statistical report of the inter-laboratory test is given in [Annex F](#).

13 Test report

The test report shall include the following items:

- a) specimen identification;
- b) the kind(s) of X-ray source used, e.g. rotating-anode W-tube;
- c) the excitation X-rays used, e.g. W-L_{II}-M_{IV};
- d) the voltage applied to the X-ray source, e.g. 30 kV;
- e) the current applied to the X-ray source, e.g. 200 mA;
- f) the glancing angle used, e.g. 1,8 mrad (0,10°);
- g) the integration time, e.g. 500 s;

- h) the method used to prepare the RM, e.g. SC1 dip method (see [Annex A](#));
- i) the atomic surface density of the RM element (Ni or Fe), e.g. Ni $1,05 \times 10^{12}$ atoms/cm²;
- j) the measurement location on the test specimen, e.g. centre of wafer;
- k) the calibration method used, i.e. procedure a) or b) in [10.2.4](#);
- l) the name(s) of the element(s) on the test specimen and the atomic surface density of each.

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Annex A (informative)

Reference materials

A.1 Reference materials for this International Standard can be prepared by the SC1 dip method [which uses standard cleaning solution 1 (a silicon wafer cleaning solution consisting of water, ammonia, and hydrogen peroxide)] or by the spin-coating method to give a surface Ni or Fe content of ca. 10^{12} atoms/cm² to 1×10^{13} atoms/cm² (see [Annex C](#)).

A.2 The Fe RM prepared by the SC1 dip method is preferred when the apparatus is installed in a higher-class cleanroom.

A.3 The Ni RM is more commonly used for routine measurements.

A.4 One or more RMs should be prepared from each lot or batch.

A.5 Specimens from the same lot or batch are assumed to have the same atomic surface density.

A.6 The calibration of RMs is discussed in [Annex C](#).

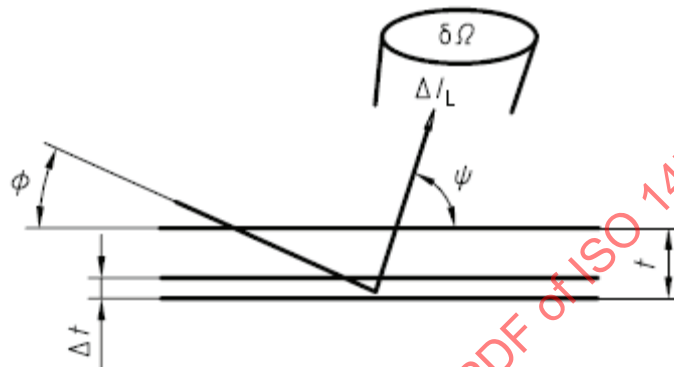
A.7 In the case of VPD-TXRF measurement, an RM consisting of a microdroplet residue containing known amounts of impurities and deposited on a hydrophobic polished or epitaxial wafer can be used.

A.8 The VPD collection efficiency is assumed to be approximately 100 %.

Annex B (informative)

Relative sensitivity factor

B.1 The relative sensitivity factor can be calculated from the model shown in [Figure B.1](#).



Key

t measurement depth
 Ω solid angle

Figure B.1 — Schematic illustration of TXRF

The X-ray intensity ΔI_L of measured spectrum L of element A, generated from the very thin layer Δt is given by:

$$\Delta I_L = I_{0,\lambda p} \exp \left\{ - \left[(\mu / \rho)_{M,\lambda p} \rho t \operatorname{cosec} \phi \right] \right\} (\mu / \rho)_{M,\lambda p} \rho \Delta t \operatorname{cosec} \phi \frac{w_A (\mu / \rho)_{A,\lambda p}}{(\mu / \rho)_{M,\lambda p}} \times \frac{\gamma_A - 1}{\gamma_A} \omega_A g_L \frac{\delta \Omega}{4\pi} \exp \left\{ - \left[(\mu / \rho)_{M,\lambda L} \rho t \operatorname{cosec} \psi \right] \right\} \quad (\text{B.1})$$

where

- $I_{0,\lambda p}$ is the incident X-ray intensity at wavelength λp ;
- $(\mu/\rho)_{M,\lambda p}$ is the mass absorption coefficient for the X-rays incident on specimen M; [2]
- ρ is the density of specimen M;
- w_A is the mass fraction of element A in the specimen;
- $(\mu/\rho)_{A,\lambda p}$ is the mass absorption coefficient for the X-rays incident on element A;
- γ_A is the jump ratio of the series shell of element A at the absorption edge; [2]
- ω_A is the fluorescence yield of the series shell of element A; [2] [3] [4]
- g_L is the relative transition probability of the measured spectrum L; [5]
- $(\mu/\rho)_{M,\lambda L}$ is the mass absorption coefficient of measured spectrum L at wavelength λ_L for specimen M.

Assuming that the value of the depth t is very small:

$$\exp \left\{ - \left[(\mu / \rho)_{M,\lambda p} \rho t \operatorname{cosec} \phi \right] \right\} = 1$$

$$\exp \left\{ - \left[(\mu / \rho)_{M,\lambda L} \rho t \operatorname{cosec} \psi \right] \right\} = 1$$

Adding double excitation of total reflection, Formula (B.1) can be expressed as follows:

$$I_L = 2 I_{0,\lambda p} w_A (\mu / \rho)_{A,\lambda p} \frac{\gamma_A - 1}{\gamma_A} \omega_A g_L \frac{\delta \Omega}{4\pi} \rho t \operatorname{cosec} \phi \quad (\text{B.2})$$

As the value of the depth t is very small:

$$w_A r t = C_A \left(A_{r,A} / N_A \right) \quad (\text{B.3})$$

where

- C_A is the atomic surface density of element A;
- $A_{r,A}$ is the atomic mass of element A;
- N_A is Avogadro's number.

From Formulae (B.2) and (B.3):

$$S_A = \frac{I_L}{C_A} = 2I_{0,\lambda p} \frac{\delta\Omega}{4\pi} (A_{r,A} / N_A) \operatorname{cosec}\phi (\mu / \rho)_{A,\lambda p} \frac{\gamma_A - 1}{\gamma_A} \omega_A g_L$$

The relative sensitivity factor, S_R is thus given by:

$$S_R = \frac{S_A}{S_{RM}} = \frac{(\mu / \rho)_{A,\lambda p} \frac{\gamma_A - 1}{\gamma_A} \omega_A g_L \cdot A_{r,A} \cdot E_A}{(\mu / \rho)_{RM,\lambda p} \frac{\gamma_{RM} - 1}{\gamma_{RM}} \omega_{RM} g_{\lambda RM} \cdot A_{r,RM} \cdot E_{RM}} \quad (\text{B.4})$$

where

$A_{r,RM}$ is the relative atomic mass of the RM element;

E_A, E_{RM} are the attenuation factors in the solid-state detector for wavelengths λ_L and λ_{RM} , respectively.

The above formula is based on the assumption that the specimen has a uniform density and a smooth surface, that monochromatized X-rays with no divergency are used, and that no multiple scattering or excitation by other elements present occurs.

B.2 When these parameters are unknown or only approximate values are required, [Table B.1](#) or [Table B.2](#) is used, provided that the RM element is Ni.

B.3 [Table B.1](#) lists calculated values of S_R , using W-L_{II}-M_{IV} as the incident X-rays and a solid-state detector with a 12,5-μm-thick Be window. [Table B.2](#) lists the calculated values of S_R using Mo-K-L_{II,III} as the incident X-rays and a solid-state detector with a 12,5-μm-thick Be window. Under different conditions (e.g. Au anode or carbon filter), the values will have to be calibrated from Formula (B.4).

Table B.1 — Relative sensitivity factors of elements for W-L_{II}-M_{IV}

Atomic no.	16	17	18	19	20	21	22
Spectrum	S-K-L _{II,III}	Cl-K-L _{II,III}	Ar-K-L _{II,III}	K-K-L _{II,III}	Ca-K-L _{II,III}	Sc-K-L _{II,III}	Ti-K-L _{II,III}
S_R	0,022 5	0,036 9	0,056 3	0,082 6	0,117	0,168	0,225
Atomic no.	23	24	25	26	27	28	29
Spectrum	V-K-L _{II,III}	Cr-K-L _{II,III}	Mn-K-L _{II,III}	Fe-K-L _{II,III}	Co-K-L _{II,III}	Ni-K-L _{II,III}	Cu-K-L _{II,III}
S_R	0,302	0,402	0,517	0,659	0,850	1,000	1,237
Atomic no.	30	33	42	46	47	50	
Spectrum	Zn-K-L _{II,III}	As-L _{III} -M _{IV,V}	Mo-L _{III} -M _{IV,V}	Pd-L _{III} -M _{IV,V}	Ag-L _{III} -M _{IV,V}	Sn-L _{III} -M _{IV,V}	
S_R	1,512	0,004 50	0,044 0	0,081 9	0,093 0	0,146	

Table B.2 — List of relative sensitivity factor of elements for Mo-K-L_{II,III}

Atomic no.	16	17	18	19	20	21	22
Spectrum	S-K-L _{II,III}	Cl-K-L _{II,III}	Ar-K-L _{II,III}	K-K-L _{II,III}	Ca-K-L _{II,III}	Sc-K-L _{II,III}	Ti-K-L _{II,III}
S_R	0,019 4	0,032 1	0,048 8	0,073 1	0,105	0,153	0,207
Atomic no.	23	24	25	26	27	28	29
Spectrum	V-K-L _{II,III}	Cr-K-L _{II,III}	Mn-K-L _{II,III}	Fe-K-L _{II,III}	Co-K-L _{II,III}	Ni-K-L _{II,III}	Cu-K-L _{II,III}
S_R	0,276	0,373	0,483	0,641	0,813	1,000	1,242
Atomic no.	30	33	42	46	47	50	73
Spectrum	Zn-K-L _{II,III}	As-K-L _{II,III}	Mo-L _{III} -M _{IV,V}	Pd-L _{III} -M _{IV,V}	Ag-L _{III} -M _{IV,V}	Sn-L _{III} -M _{IV,V}	Ta-L _{III} -M _{IV,V}

Table B.2 (continued)

S_R	1,538	2,445	0,041 2	0,077 7	0,089 4	0,142	2,253
Atomic no.	74	78	79	80	82	92	
Spectrum	W-L _{III} -M _{IV,V}	Pt-L _{III} -M _{IV,V}	Au-L _{III} -M _{IV,V}	Hg-L _{III} -M _{IV,V}	Pb-L _{III} -M _{IV,V}	U-L _{III} -M _{IV,V}	
S_R	2,566	3,643	3,734	4,015	5,023	5,574	

B.4 Correction factors can be made to allow for the instrumental conditions.

B.5 Instead of using relative sensitivity factors, a calibration curve can be prepared using calibration specimens.

B.6 If such a calibration curve is used, recalibration with RMs will be necessary at appropriate intervals.

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Annex C (informative)

Preparation of reference materials^[6]

C.1 Requirements on RMs for TXRF

TXRF is a highly sensitive method of analysis, and the analyte content of RMs is very low compared with that for other analytical methods. Even a little contamination will make an RM unsuitable. Thus, RMs are not suitable for permanent use. Fresh RMs have to be prepared and calibrated from time to time. It is therefore preferable to use an RM which can be prepared and calibrated as easily as possible.

The area of incidence and the flux density of the excitation X-rays vary widely with the glancing angle, and the intensity of the detector signal depends on the relative disposition of the specimen and the X-ray detector.^[7] When an RM is calibrated, there is no easy quantitative analytical method that can determine the analyte content exclusively in the small area on the specimen where the TXRF analysis takes place. Therefore, for the preparation of a reliable working RM, it is desirable to use a method which deposits the reference element uniformly over the whole surface of the wafer.

In addition, long-term quality stability is required.

Specific methods of preparing an RM which meet the above-mentioned requirements are the SC1 dip method [also known as the IAP (immersion in alkaline hydrogen peroxide solution) method]^{[8][9]} and the spin-coating method.^[10] The latter method, however, tends to cause particle formation. Another possible method is the microdrop method^[7] by which a certain quantity of the RM element is deposited on a part of the surface, though the resultant specimen is not uniform.

C.2 Silicon wafer for the RM

The surface density of the contamination on the silicon wafer used as the RM is required to be less than 10^{10} atoms/cm². Wafers used for 16M or higher integration DRAM (dynamic random-access memory) are suitable.

If the crystallographic orientation of the wafer varies, the conditions for the generation of spectral-interference peaks will vary.^[11] Hence the azimuth of the chemomechanically polished or epitaxial wafer to be used should preferably be identical with that of the wafer under test.

C.3 The RM element

Transition metals with a high generation efficiency are appropriate for use as the RM element. Of such elements, Ni and Fe are recommended for the following reasons:

- a) If the W-L_{II}-M_{IV} line is used as the X-ray source, there will be some overlap of the Cu peak with the escape peak of the scattered W-L_{II}-M_{IV} line. This can result in errors due to the deconvolution process;
- b) If the W-L_{II}-M_{IV} line is used as the X-ray source, there will be some overlap of the Zn peak with the scattered W-L_{II}-M_{IV} line and with the Compton peaks. This can result in errors, depending how the baseline is drawn;
- c) The results of the inter-laboratory test programme have shown that Fe and Ni are comparatively stable, i.e. they are deposited in the form of a uniform film and they remain within the native-oxide film;^[12]

- d) Fe-containing airborne particles are fairly common, and when they fall onto the wafer surface or adhere to the detector window they will cause the generation of spurious peaks;

More careful treatment of an Fe RM is therefore required, as compared to RMs utilizing other elements, e.g. Ni. It was observed, however, by all the participants in the inter-laboratory test programme that the Fe RM was not contaminated.

- e) It is not known how the use of Co, V, or Mn as the RM element would affect the native-oxide film. Also, the behaviour of these elements inside or outside the native-oxide film on the silicon surface is unknown, and the precision of such RMs has not been demonstrated by inter-laboratory testing.

C.4 Procedures for preparing RMs

C.4.1 SC1 dip method[8][9]

The wafer surface is first cleaned so as to obtain a surface contamination level of 10^{10} atoms/cm² or less. The wafer is then dipped in an SC1 cleaning solution (water, ammonia, and hydrogen peroxide in the ratio 5:0,1 to 1:1 by volume) containing the ions of the RM element in order to deposit a certain quantity of the RM element in the native-oxide film.

It should be noted, however, that it is difficult to deposit certain elements, e.g. Ni, uniformly over the whole range of concentration of ammonia in the solution. It is therefore recommended that a 5:0,25:1 solution be used, as this provides satisfactorily uniform surface deposition.

The use of SC1 cleaning solution has the advantage that etching and formation of the native-oxide film proceed concurrently, thereby maintaining the thickness of the native-oxide film whilst allowing the impurity element in the solution to be taken up by the native-oxide film.

C.4.2 Spin-coating method[10]

The wafer surface is first cleaned so as to obtain a hydrophobic surface with a contamination level of 10^{10} atoms/cm² or less. A droplet of an acidic solution containing the ions of the RM element is then dispersed over the entire wafer surface. The wafer is held in this state for a certain period of time to allow the RM element to be dispersed on the silicon substrate and is then spun dry.

This method provides satisfactory uniformity of the surface contaminant. More than one element can be deposited at the same time.

C.4.3 Microdrop method[11]

The wafer surface is first cleaned so as to obtain a hydrophobic surface with a contamination level of 10^{10} atoms/cm² or less. A droplet of known volume of a solution containing the ions of the RM element is then deposited on the wafer surface and allowed to dry.

The surface density of the deposited element depends on its concentration in the solution and the volume of the droplet. It should be noted, however, that the intensity of the fluorescence X-rays can vary considerably depending upon the relative positions of the droplet residue and the detector.[7] The diameter of the droplet residue will also affect the intensity[12] (see [Annex E](#)).

C.5 Calibration of RMs

RMs are calibrated as soon as possible after preparation.

Wafers prepared by the SC1 or the spin-coating method are calibrated by AAS, ICP-MS or some other method relevant to the RM element contained in the native-oxide film on the wafer and recovered by the VPD specimen preparation method (see [Annex D](#)).

If hydrofluoric acid is used to gather the impurities, including Ni, from the oxide film, the impurities present in the acid may be a source of difficulty. Often, therefore, the native-oxide film on the wafer

surface is dissolved using vapourized hydrofluoric acid to recover the RM element. Radioactivation analysis is also used, which avoids circumferential contamination after radioactivation, such as occurs during VPD specimen preparation and the subsequent analysis.

Rutherford backscattering and forward-scattering methods, including the method known as heavy-ion backscattering spectroscopy, can be used to calibrate RMs directly.

RMs prepared by the microdrop method do not need to be calibrated as the surface density of the deposited element can be calculated from its concentration in the solution and the volume of the droplet. A calculation procedure which gives reliable results is given in SEMI Standard M33.

C.6 Recommended RMs

The SC1 method produces RM wafers having contaminant-density levels of 10^{11} atoms/cm² to 10^{13} atoms/cm² with good surface-density and depth uniformity. The repeatability of this method is such that up to 25 practically identical specimens can be produced at the same time.

In the case of the spin-coating method, the depth distribution of the deposited element is not constant. Though this would lead to an error in the analytical results, the international inter-laboratory test programme demonstrated that specimens prepared by the spin-coating method gave satisfactory reproducibility.

The predominant problem associated with the microdrop method is that the calculation procedure for converting the solution concentration into a surface density is not yet reliably established. Also, any error in setting the coordinates would lead to a difference in the fluorescence X-ray intensity. Further errors may be introduced by the fact that the distribution of the elements on the RM and the test specimen are always different. This method is simple, however, and applicable to lower surface densities.

It is recommended that the SC1 method and the spin-coating method be used.

C.7 Storage and lifetime of RMs

RMs should be stored in the same container as the blank RM and in the same environment as the TXRF instrument.

Despite these precautions, airborne particles can be deposited on RMs during measurement.

It is probable that the native-oxide film grows and hydrocarbons are deposited on RMs during storage in the container.

It is reasonable to assess the level of contamination of the RM and the blank RM from the apparent level of contamination on the blank RM after only making a background correction. If the apparent level of contamination is greater than 10 % of the surface density of the element on the RM, the RM and the blank RM are considered to have exceeded their useful life and should be replaced. The normal life expectancy of RMs is less than 6 mo.

C.8 Caution

The user should be aware that silicon surfaces can be covered by both particulate-type and film-type contamination, and that some impurity elements, e.g. Zn, can occur in both of these forms. The fluorescence X-ray intensity depends upon the physical state and the depth of the contamination. The elemental levels measured for particulate-type contamination and for the residue produced by a VPD scanning droplet will be higher than the actually present at the analysed sample surface if film-type RMs are used for quantification purposes. This is, however, the most reliable means of quantifying levels of contamination on unknown specimens.

Annex D (informative)

VPD-TXRF method

D.1 General information

The standard TXRF method is valid in the range from 10^{10} atoms/cm² to 10^{13} atoms/cm². The practical measurement range can be improved upon by the use of VPD-TXRF, with which the detection range is 5×10^8 atoms/cm² to 5×10^{12} atoms/cm². In the VPD-TXRF method, the contaminants are collected from the whole wafer surface and deposited in the form of a small residue which is then analysed by TXRF.

D.2 Collection of contamination elements

The contamination elements are collected by decomposing the native-oxide film present on the wafer surface. This can be achieved either by exposing the wafer surface to HF vapour in a VPD chamber or by scanning an HF droplet over the surface of a polished or epitaxial silicon wafer specimen.

An HF-treated surface becomes hydrophobic. Thus, any droplets on the wafer surface will tend to become spherical as the contact angle between the droplets and wafer surface exceeds 1,05 rad (60°), so these droplets will gather into one droplet. Alternatively, by sandwiching one drop of HF or HF + HNO₃ between the container and wafer it is possible to expose the whole wafer surface simultaneously.^[13] The chemicals should be of ultra-pure grade and the environment in which this operation is carried out should be ISO Class 4 or better. The droplet obtained can also be analysed by the AAS or the ICP-MS method.

NOTE Patents concerning methods of collecting contamination elements have been applied for in many countries. ISO cannot give any authoritative or comprehensive information about the existence, validity, or scope of such patent rights. Users of this International Standard are expressly advised that determination of any such patent rights and the risk of infringement of such rights are entirely their own responsibility.

D.3 Producing the droplet residue

The spherical droplet is dried on the silicon wafer surface from which the contamination elements have been collected or on another wafer which has been freshly made hydrophobic. The accuracy is dependent on the size (area and height) of the residue: the smaller the size, the better the accuracy. It is well known that a hydrophobic silicon wafer strongly attracts and adsorbs airborne particles or chemicals in the cleanroom. Use of a drying chamber with a vacuum system or a flow of clean, inert gas at ambient temperature is recommended.

D.4 Measurement by TXRF

D.4.1 Size of residue

The residue is small. The artificial residue employed in the inter-laboratory test programme had a diameter of 350 µm and a height of 50 nm.

D.4.2 Finding the residue spot

It is very difficult to find the residue spot unless the precise position is recorded, the quick-search system of the TXRF instrument is used or a spot-mark is made with nitric acid.

D.4.3 Effect of residue size

When the residue is not small, the measurement error becomes greater. In addition, salts of different metallic elements often become separated and the distribution of such elements is not homogeneous. This inhomogeneity makes the determination of multi-element contamination less accurate.

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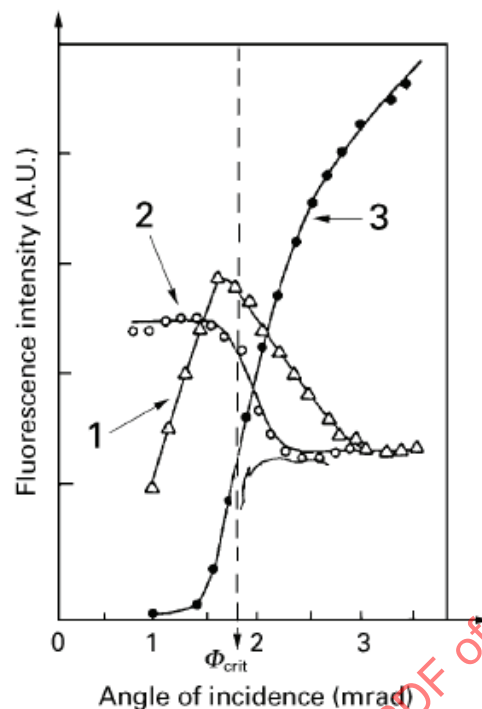
Annex E (informative)

Glancing-angle settings

E.1 TXRF anglescans of ideal specimens^[14]

TXRF anglescans using an Mo-K-L_{II,III} X-ray source are shown in [Figure E.1](#) for an ideal specimen consisting of a polished silicon wafer. Of particular importance is the difference in the anglescans of the particulate-type and film-type contamination, although the atomic surface density is the same in each case. At about 1 mrad (0,06°), there is a very large difference between the TXRF signals for the particulate-type and film-type contamination for the same surface density. At about 1,3 mrad (0,07°), there is no difference between the two signals. This angle is sometimes called the “crossover angle” because it is the angle at which the anglescans for the two types of contamination cross each other.

The importance of this crossover angle is clear when we consider the anglescan of an RM and that of an unknown specimen. Assume for the moment that the RM has an anglescan of the film-type form in [Figure E.1](#). Now if the unknown specimen has an anglescan similar to the film-type curve (plated or sputtered) of [Figure E.1](#), any angle can be used for the quantification. However, if the unknown has an anglescan similar to the particulate-type curve (residue) in [Figure E.1](#), then only the measurement at the crossover angle gives correct quantification. If in this case the measurement is made instead at a low angle, such as 1 mrad (56 % of the theoretical critical angle), the accuracy of the quantification is in error by the ratio of the signal intensities shown in [Figure E.1](#) at 1 mrad. In this figure, the error would be about 2,5 times. If an even lower angle were used, the error could reach five times or more.



Key

- 1 plated
- 2 residue
- 3 bulk

Figure E.1 — Experimental curves showing the angular dependence of the fluorescence

In actual analysis, it is difficult to check the anglescan of each unknown, because of the very low signal intensities at low contamination levels. Therefore, one cannot always know whether or not there is a major error in the quantification if a lower angle is used.

The only angle which ensures accurate quantification in the measurement of ideal specimens is the crossover angle, and it is therefore this angle which is recommended. In real RMs, the anglescan is sometimes not exactly like the film-type anglescan in [Figure E.1](#), but is somewhat broader in width. In this case, the crossover point becomes a range, perhaps between 1,2 mrad (67 % of the theoretical critical angle) and 1,5 mrad (84 % of the theoretical critical angle).

E.2 Anglescan of film-type contamination[6][15]

The depth of measurement is defined as the penetration depth at which the X-ray intensity decreases by 1/e. In general, this depth varies with the energy of the X-rays, the glancing angle and the matrix electron density. For the TXRF analysis of silicon surface types covered by this International Standard and the recommended range of the glancing angles, the calculated penetration depth values are less than 5 nm. The intensity of the fluorescence X-rays from each contamination element varies with the glancing angle. [Figure E.2](#) shows the results of anglescans, using an Ni-K-L_{II,III} X-ray source, of three Ni RMs made by the SC1 dip method: Ni-1 with ca. 10¹¹ atoms/cm², Ni-2 with ca. 10¹² atoms/cm² and Ni-3 with ca. 10¹⁴ atoms/cm². The anglescan curves are normalized with respect to their maximum value. The curves show the differences between the deposited-film types. Ni-3 is assumed to be particle-forming NiSi precipitates. [Figure E.3](#) shows the results of anglescans, using an Ni-K-L_{II,III} X-ray source, of Ni-3 by three laboratories. The anglescan curves are the same in each case.