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Standard Test Methods for Estimating the Depth of Decarburization of Steel Specimens¹

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This standard has been approved for use by agencies of the U.S. Department of Defense.

INTRODUCTION

These test methods may be used to estimate the average or greatest depth of decarburization in hardened or non-hardened steel products. The test methods described range from simple screening tests to more statistically rigorous test methods depending upon the needs of the investigation.

1. Scope

1.1 These test methods cover procedures for estimating the depth of decarburization of steels irrespective of the composition, matrix microstructure, or section shape. The following basic procedures may be used:

- 1.1.1 Screening methods.
- 1.1.2 Microscopical methods.
- 1.1.3 Microindentation hardness methods.
- 1.1.4 Chemical analysis methods.

1.2 In case of a dispute, the rigorous quantitative or lineal analysis method (see 7.3.5 and 7.3.6) shall be the referee method. These methods can be employed with any cross-sectional shape. The chemical analytical methods generally reveal a greater depth of decarburization than the microscopical methods but are limited to certain simple shapes and by availability of equipment. These techniques are generally reserved for research studies. The microindentation hardness method is suitable for accurate measurements of hardened structures with relatively homogeneous microstructures.

1.3 The values stated in SI units are to be regarded as standard. No other units of measurement are included in this standard.

1.4 *This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety, health, and environmental practices and determine the applicability of regulatory limitations prior to use.*

¹ These test methods are under the jurisdiction of ASTM Committee E04 on Metallography and are the direct responsibility of Subcommittee E04.14 on Quantitative Metallography.

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1.5 *This international standard was developed in accordance with internationally recognized principles on standardization established in the Decision on Principles for the Development of International Standards, Guides and Recommendations issued by the World Trade Organization Technical Barriers to Trade (TBT) Committee.*

2. Referenced Documents

2.1 *ASTM Standards:*²

- A941 Terminology Relating to Steel, Stainless Steel, Related Alloys, and Ferroalloys
- E3 Guide for Preparation of Metallographic Specimens
- E7 Terminology Relating to Metallography
- E340 Practice for Macroetching Metals and Alloys
- E350 Test Methods for Chemical Analysis of Carbon Steel, Low-Alloy Steel, Silicon Electrical Steel, Ingot Iron, and Wrought Iron
- E384 Test Method for Microindentation Hardness of Materials
- E407 Practice for Microetching Metals and Alloys
- E415 Test Method for Analysis of Carbon and Low-Alloy Steel by Spark Atomic Emission Spectrometry
- E1951 Guide for Calibrating Reticles and Light Microscope Magnifications

3. Terminology

3.1 *Definitions:*

3.1.1 For definitions of terms used in these test methods, see Terminology E7 and Terminology A941.

3.2 *Definitions of Terms Specific to This Standard:*

² For referenced ASTM standards, visit the ASTM website, www.astm.org, or contact ASTM Customer Service at service@astm.org. For *Annual Book of ASTM Standards* volume information, refer to the standard's Document Summary page on the ASTM website.

3.2.1 *average depth of decarburization*—the mean value of five or more measurements of the total depth of decarburization.

3.2.2 *average free-ferrite depth*—the mean value of five or more measurements of the depth of complete decarburization.

3.2.3 *complete decarburization*—loss of carbon content at the surface of a steel specimen to a level below the solubility limit of carbon in ferrite so that only ferrite is present.

3.2.4 *free-ferrite depth*—the perpendicular distance from the surface of the specimen to that location where the structure is no longer fully ferritic, that is, other transformation products are observed.

NOTE 1—The term *free ferrite* has also been used to describe globular, isolated grains of proeutectoid ferrite in the microstructure of medium-carbon hypoeutectoid steels.

3.2.5 *maximum depth of decarburization*—the largest measured value of the total depth of decarburization.

3.2.6 *partial decarburization*—loss of carbon content at the surface of a steel specimen to a level less than the bulk carbon content of the unaffected interior but greater than the room temperature solubility limit of carbon in ferrite.

3.2.7 *total depth of decarburization*—the perpendicular distance from the specimen surface to that location in the interior where the bulk carbon content is reached; that is, the sum of the depths of complete and partial decarburization.

4. Summary of Test Method

4.1 These test methods are designed to detect changes in the microstructure, hardness, or carbon content at the surface of steel sections due to decarburization. The depth of decarburization is determined as the depth where a uniform microstructure, hardness, or carbon content, typical of the interior of the specimen, is observed.

5. Significance and Use

5.1 These test methods are used to detect surface losses in carbon content due to heating at elevated temperatures, as in hot working or heat treatment.

5.2 Results of such tests may be used to qualify material for shipment according to agreed upon guidelines between purchaser and manufacturer, for guidance as to machining allowances, or to assess the influence of processing upon decarburization tendency.

5.3 Screening tests are simple, fast, low-cost tests designed to separate non-decarburized samples from those with appreciable decarburization. Based on the results of such tests, the other procedures may be utilized as applicable.

5.4 Microscopical tests require a metallographically polished cross section to permit reasonably accurate determination of the depth and nature of the decarburization present. Several methods may be employed for estimation of the depth of decarburization. The statistical accuracy of each varies with the amount of effort expended.

5.5 Microindentation hardness methods are employed on polished cross sections and are most suitable for hardened specimens with reasonably uniform microstructures. This pro-

cedure can be used to define the depth to a specific minimum hardness or the depth to a uniform hardness.

5.6 Chemical analytical methods are limited to specimens with simple, uniform shapes and are based on analysis of incremental turnings or after milling at fixed increments.

5.7 Microscopical tests are generally satisfactory for determining the suitability of material for intended use, specification acceptance, manufacturing control, development, or research.

6. Sampling

6.1 Samples should be taken at locations that are representative of the bulk specimen. The location and number of samples taken depends on the nature of the material to be tested and will be defined upon agreements between manufacturer and purchaser.

6.2 Specimens for screening tests using bulk hardness tests, such as the Rockwell test, should be small enough so that they can be properly supported on the anvil of the tester. The specimen surface should not be altered except for scale removal (if present) using a method that will not alter the subsurface metal.

6.3 Specimens for the microscopical methods or for micro-indentation hardness tests or for macroscopic screening methods should be cut from the bulk specimen perpendicular to the longitudinal axis of the product so that measurements are made on a transverse plane. This procedure permits determination of the variation of decarburization around the periphery of the specimen.

6.3.1 For specimens up to about 2.5-cm diameter, the entire cross section is polished and examined. For larger cross sections, one or more specimens shall be prepared to assess variations in surface decarburization. Figs. 1-3 show examples of typical sampling schemes that may be used for larger sections; the sampling scheme for large sections should be determined upon mutual agreement between manufacturer and purchaser.

6.4 Specimens for chemical analytical methods must be of sufficient length so that the weight of incremental turnings is adequate for chemical analysis or the size of milled surfaces is large enough for sparking yet small enough to fit in the specimen holder.

7. Procedure

7.1 Screening Methods:

7.1.1 *Bulk Surface Hardness*—For hardened specimens, particularly those in the as-quenched condition, a short section of the material to be heat treated is cut and heat treated in the same manner, or along with, the material of interest. The test specimen, however is not tempered. Any scale on the test piece is removed by wire brushing, glass-bead blasting, etc., and hardness tested, usually with the Rockwell C scale. The presence of decarburization is indicated by the difference between the surface hardness and the theoretical maximum hardness for the carbon content of the steel. This method is most suitable for those steels with bulk carbon contents below about 0.55 % carbon but will detect gross decarburization in steels with higher bulk carbon contents. The method is not