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# Heat Treatment and Properties of Iron and Steel

Thomas G. Digges, Samuel J. Rosenberg, and Glenn W. Geil

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## National Bureau of Standards Monograph 88

Issued November 1, 1966 Supersedes Circular 495 and Monograph 18

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Library of Congress Catalog Card No. 66-61523

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## Heat Treatment and Properties of Iron and Steel

Thomas G. Digges,<sup>1</sup> Samuel J. Rosenberg,<sup>1</sup> and Glenn W. Geil

This Monograph is a revision of the previous NBS Monograph 18. Its purpose is to provide an understanding of the heat treatment of iron and steels, principally to those unacquainted with this subject. The basic principles involved in the heat treatment of these materials are presented in simplified form. General heat treatment procedures are given for annealing, normalizing, hardening, tempering, case hardening, surface hardening, and special treatments such as austempering, ausforming, martempering and cold treatment. Chemical compositions, heat treatments, and some properties and uses are presented for structural steels, tool steels, stainless and heatresisting steels, precipitation-hardenable stainless steels and nickel-maraging steels.

#### 1. Introduction

The National Bureau of Standards receives many requests for general information concerning the heat treatment of iron and steel and for directions and explanations of such processes. This Monograph has been prepared to answer such inquiries and to give in simplified form a working knowledge of the basic theoretical and practical principles involved in the heat treatment of iron and steel. The effects of various treatments on the structures and mechanical properties of these materials are described. Many theoretical aspects are discussed only briefly or omitted entirely, and in some instances, technical details have been neglected for simplicity. The present Monograph supersedes Circular 495, which was published in 1950, and Monograph 18 (1960).

Heat treatment may be defined as an operation or combination of operations that involves the heating and cooling of a solid metal or alloy for the purpose of obtaining certain desirable conditions or properties. It is usually desired to preserve, as nearly as possible, the form, dimensions, and surface of the piece being treated.

Steels and cast irons are essentially alloys of iron and carbon, modified by the presence of other elements. Steel may be defined as an alloy of iron and carbon (with or without other alloying elements) containing less than about 2.0 percent of carbon, usefully malleable or forgeable as initially cast. Cast iron may be defined as an alloy of iron and carbon (with or without other alloying elements) containing more than 2.0 percent of carbon, not usually malleable or forgeable as initially cast. For reasons that will be apparent later, the dividing line between steels and cast irons is taken at 2.0 percent of carbon, even though certain special steels contain carbon in excess of this amount. In addition to carbon, four other elements are normally present in steels and in cast irons. These are manganese, silicon, phosphorus, and sulfur.

Steels may be broadly classified into two types, (1) carbon and (2) alloy. Carbon steels owe their properties chiefly to the carbon. They are frequently called straight or plain carbon steels. Alloy steels are those to which one or more alloying elements are added in sufficient amounts to modify certain properties. The properties of cast iron also may be modified by the presence of alloying elements—such irons are called alloy cast irons.

#### 2. Properties of Iron

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Since iron is the basic element of steel, a knowledge of some of its properties is a prerequisite to an understanding of the fundamental principles underlying the heat treatment of steels.

#### 2.1. Transformation Temperatures

If a molten sample of pure iron were allowed to cool slowly and the temperature of the iron were measured at regular intervals, an ideal-

<sup>1</sup> Retired.

ized (equilibrium) time-temperature plot of the data would appear as shown in figure 1. The discontinuities (temperature arrests) in this curve are caused by physical changes in the iron.

The first arrest at 2,800 °F marks the temperature at which the iron freezes. The other arrests (known as transformation temperatures or critical points) mark temperatures at which certain internal changes take place in the solid iron. Some of these temperatures are very important in the heat treatment of steel.

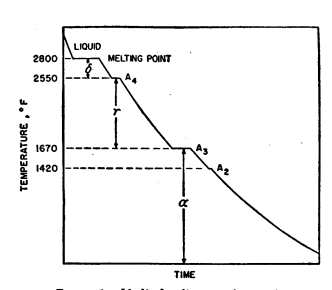
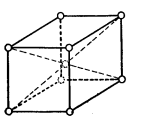


FIGURE 1. Idealized cooling curve for pure iron.

The atoms in all solid metals are arranged in some definite geometric (or crystallographic) pattern. The atoms in iron, immediately after freezing, are arranged in what is termed the body-centered cubic system. In this crystal structure the unit cell consists of a cube with an iron atom at each of the eight corners and another in the center (fig. 2, a). Each of the many individual grains (crystals) of which the solid metal is composed is built up of a very large number of these unit cells, all oriented alike in the same grain. This high-temperature form of iron is known as delta ( $\delta$ ) iron.

At 2,550 °F (the A<sub>4</sub> point), iron undergoes; an allotropic transformation (fig. 1); that is, the arrangement of the atoms in the crystal changes. The new crystal structure is facecentered cubic, and the unit cell again consists, of a cube with an iron atom at each of the eight corners, but with an iron atom in the center of each of the six faces instead of one in the center of the cube (fig. 2, b). This form is known as gamma ( $\gamma$ ) iron. At 1,670 °F (the A<sub>3</sub> point) iron undergoes another allotropic



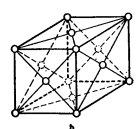


FIGURE 2. Crystal structure of iron. (a) Body-centered cubic (alpha and delta iron); (b) Face-centered cubic (gamma iron).

transformation and reverts to the bodycentered cubic system (fig. 2, a). This structure, which is crystallographically the same as delta iron, is stable at all temperatures below the  $A_3$  point and is known as alpha ( $\alpha$ ) iron (fig. 5, A). The arrest at 1,420 °F (known as the  $A_2$  point) is not caused by an allotropic change, that is, a change in crystal structure. It marks the temperature at which iron becomes ferromagnetic and is therefore termed the magnetic transition. Above this temperature iron is nonmagnetic.

These various temperature arrests on cooling are caused by evolutions of heat. On heating, the arrests occur in reverse order and are caused by absorption of heat. The critical points may be detected also by sudden changes in other physical properties, for instance, expansivity or electrical conductivity.

#### 2.2. Mechanical Properties

Iron is relatively soft, weak<sup>\*</sup>, and ductile and cannot be appreciably hardened by heat treatment. Its tensile strength at room temperature is about 40,000 lb/in.<sup>2</sup>, its yield strength is about 20,000 lb/in.<sup>2</sup>, and its Brinell hardness is about 80. The modulus of elasticity is about 29,000,000 lb/in.<sup>2</sup>. The strength and hardness can be increased, with corresponding decrease in ductility, by cold working.

### 3. Alloys of Iron and Carbon

The properties of iron are affected very markedly by additions of carbon. It should be realized that in discussing iron-carbon alloys, we actually are dealing with plain carbon steels and cast irons.

#### 3.1. Iron-Carbon Phase Diagram

The complete iron-carbon phase (or constitutional) diagram represents the relationship between temperatures, compositions, and structures of all phases that may be formed by iron and carbon under conditions of equilibrium (very slow cooling). A portion of this diagram for alloys ranging up to 6.7 percent of carbon is reproduced in figure 3; the upper limit of carbon in cast iron is usually not in excess of 5 percent. The lefthand boundary of the diagram represents pure iron, and the right-hand boundary represents the compound iron carbide,  $Fe_3C$ , commonly called cementite.

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<sup>\*</sup> Iron grown as single-crystal whiskers may exhibit astoundingly high strengths.

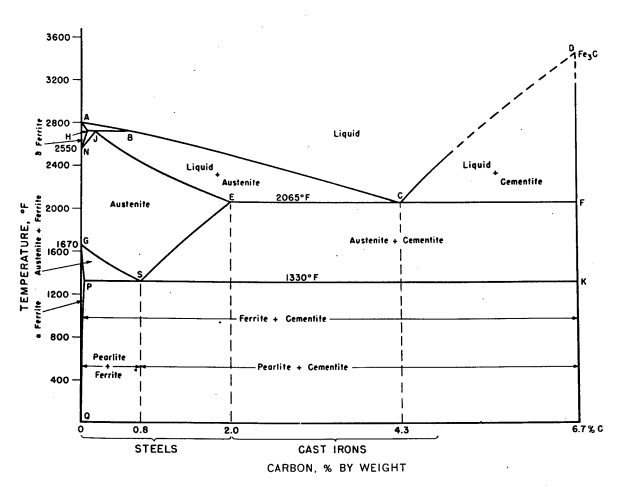


FIGURE 3. Iron-carbon phase diagram.

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The beginning of freezing of the various iron-carbon alloys is given by the curve ABCD. termed the liquidus curve. The ending of freezing is given by the curve AHJECF, termed the solidus curve. The freezing point of iron is lowered by the addition of carbon (up to 4.3%) and the resultant alloys freeze over a range in temperature instead of at a constant temperature as does the pure metal The alloy containing 4.3 percent of cariron. bon, called the eutectic alloy of iron and cementite, freezes at a constant temperature as indicated by the point C. This temperature is 2,065 °F, considerably below the freezing point of pure iron.

Čarbon has an important effect upon the transformation temperatures (critical points) of iron. It raises the  $A_4$  temperature and lowers the  $A_3$ . This effect on the  $A_3$  temperature is very important in the heat treatment of carbon and alloy structural steels, while that on the  $A_4$  is important in the heat treatment of certain high alloy steels, particularly of the stainless types.

It is possible for solid iron to absorb or dissolve carbon, the amount being dependent upon the crystal structure of the iron and the temperature. The body-centered (alpha or delta) iron can dissolve but little carbon, whereas the face-centered (gamma) iron can dissolve a considerable amount, the maximum being about 2.0 percent at 2,065 °F (fig. 3). This solid solution of carbon in gamma iron is termed austenite. The solid solution of carbon in delta iron is termed delta ferrite, and the solid solution of carbon in alpha iron is termed alpha ferrite, or, more simply, ferrite.

The mechanism of solidification of ironcarbon alloys, especially those containing less than about 0.6 percent of carbon, is rather complicated and is of no importance in the heat treatment of carbon steels and cast irons. It is sufficient to know that all iron-carbon alloys containing less than 2.0 percent of carbon (that is, steel) will, immediately or soon after solidification is complete, consist of the single phase austenite. Cast irons will consist of two phases immediately after solidification-austenite and cementite ( $Fe_3C$ ). Under some conditions this cementite formed on cooling through the temperature horizontal ECF will decompose partly or completely into austenite and graphite (carbon).

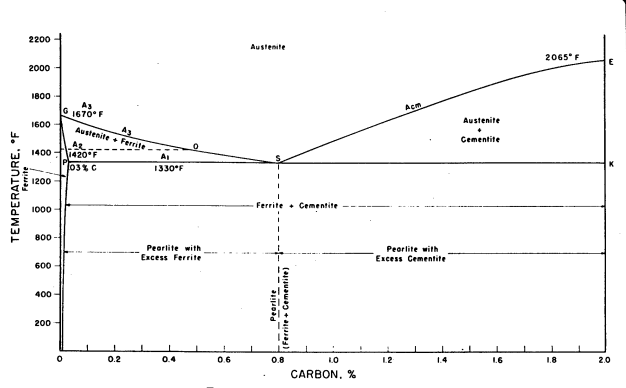


FIGURE 4. Phase diagram for carbon steels.

The part of the iron-carbon diagram that is concerned with the heat treatment of steel is reproduced on an expanded scale in figure 4. Regardless of the carbon content, steel exists as austenite above the line GOSE. Steel of composition S (0.80% of carbon) is designated as "eutectoid" steel, and those with lower or higher carbon as "hypoeutectoid" and "hypereutectoid," respectively.

A eutectoid steel, when cooled at very slow rates from temperatures within the austenitic field, undergoes no change until the tempera-ture horizontal *PSK* is reached. At this temperature (known as the A<sub>1</sub> temperature), the austenite transforms completely to an aggregate of ferrite and cementite having a typical lamellar structure (fig. 5, D and E). This aggregate is known as pearlite and the  $A_1$ temperature is, therefore, frequently referred to as the pearlite point. Since the  $A_1$  transformation involves the transformation of austenite to pearlite (which contains cementite-Fe<sub>3</sub>C), pure iron does not possess an A<sub>1</sub> transformation (fig. 4). Theoretically, iron must be alloyed with a minimum of 0.03 percent of carbon before the first minute traces of pearlite can be formed on cooling (point P, fig. 4). If the steel is held at a temperature just below  $A_1$ , (either during cooling or heating), the carbide in the pearlife tends to coalesce into globules or spheroids. This phenomenon, known as spheroidization, will be discussed subsequently.

Hypoeutectoid steels (less than 0.80% of carbon), when slowly cooled from temperatures above the  $A_3$ , begin to precipitate ferrite when the  $A_3$  line (GOS-fig. 4) is reached. As the temperature drops from the  $A_3$  to  $A_1$ , the precipitation of ferrite increases progressively and the amount of the remaining austenite decreases progressively, its carbon content being increased. At the A<sub>1</sub> temperature the remaining austenite reaches eutectoid composition (0.80% of carbon—point S, fig. 4) and, upon further cooling, transforms completely into pearlite. The microstructures of slowly cooled hypoeutectoid steels thus consist of mixtures of ferrite and pearlite (fig. 5, B and C). The lower the carbon content, the higher is the temperature at which ferrite begins to precipitate and the greater is the amount in the final structure.

Hypereutectoid steels (more than 0.80% of carbon) when slowly cooled from temperatures above the  $A_{cm}$ , begin to precipitate cementite when the  $A_{cm}$  line (SE—fig. 4) is reached. As the temperature drops from the  $A_{cm}$  to  $A_1$ , the precipitation of cementite increases progressively and the amount of the remaining austenite decreases progressively, its carbon content being depleted. At the  $A_1$  temperature the remaining austenite reaches eutectoid composition ( $0.8\bar{0}\%$  of carbon) and, upon further cooling, transforms completely into pearlite. The microstructures of slowly cooled hypereutectoid steels thus consist of mixtures of cementite and pearlite (fig. 5, F). The higher the carbon content, the higher is the temperature at which cementite begins to precipitate

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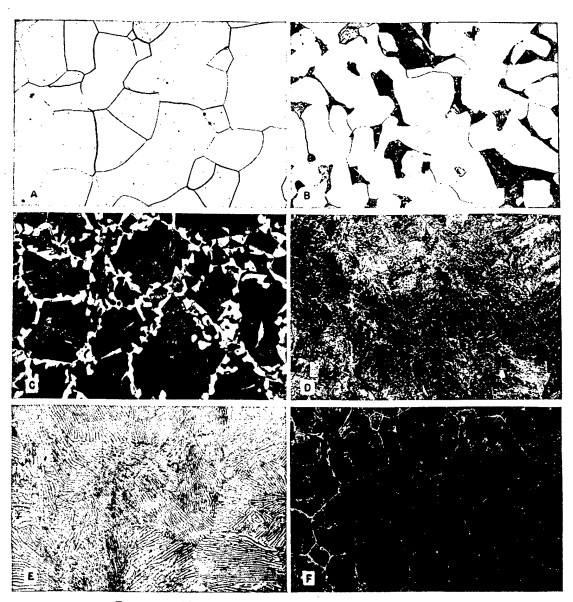


FIGURE 5. Microstructural constituents of slowly cooled carbon steels.

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A, Ferite (a iron). All grains are of the same composition.  $\times 100$ . B, 0.25% carbon. Light areas are ferrite grains. Dark areas are pearlite.  $\times 100$ . C, 0.5% carbon. Same as B but higher carbon content results in more pearlite and less ferrite.  $\times 100$ . D, 0.8% carbon. All pearlite.  $\times 100$ . E, Same as D. At higher magnification the lamellar structure of pearlite is readily observed.  $\times 500$ . F, 1.3% carbon. Arealite plus excess cementite as network.  $\times 100$ . All etched with either picral or nital.

and the greater is the amount in the final structure.

The temperature range between the  $A_1$  and A<sub>3</sub> points is called the critical or transformation range. Theoretically, the critical points in any steel should occur at about the same temperatures on either heating or cooling very slowly. Practically, however, they do not since the  $A_3$  and  $A_1$  points, affected but slightly by the rate of heating, are affected tremendously by the rate of cooling. Rapid rates of heating raise these points only slightly, but rapid rates of cooling lower the temperatures of transfor-

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mation considerably. To differentiate between the critical points on heating and cooling, the small letters "c" (for "chauffage" from the French, meaning heating) and "r" (for "refroidissement" from the French, meaning cooling) are added. The terminology of the critical points thus becomes Ac<sub>3</sub>, Ar<sub>3</sub>, Ac<sub>1</sub>, Ar<sub>1</sub>, etc. The letter "e" is used to designate the occurrence of the points under conditions of extremely slow cooling on the assumption that this represents equilibrium conditions ("e" for equilibrium); for instance, the Ae<sub>3</sub>, Ae<sub>1</sub>, and Ae<sub>cm</sub>.

## 3.2. Correlation of Mechanical Properties With Microstructures of Slowly Cooled Carbon Steels

Some mechanical properties of pearlite formed during slow cooling of a eutectoid (0.80% of carbon) steel are approximately as follows:

Tensile strength-115,000 lb/in<sup>2</sup>. Yield strength-60,000 lb/in<sup>2</sup>.

Brindell hardness number-200.

The amount of pearlite present in a slowly cooled hypoeutectoid steel is a linear function of the carbon content, varying from no pearlite, when no carbon is present (the very slight amount of carbon soluble in alpha iron may be neglected), to all pearlite at 0.80 percent of carbon. The balance of the structure of hypoeutectoid steels is composed of ferrite, the mechanical properties of which were given in a preceding section. Since the mechanical properties of aggregates of ferrite and pearlite are functions of the relative amounts of these two constituents, the mechanical properties of slowly cooled hypoeutectoid steels are also lin-ear functions of the carbon content, varying between those of iron at no carbon to those of pearlite at 0.80 percent of carbon (fig. 6).

## 4. Decomposition of Austenite

In alloys of iron and carbon, austenite is stable only at temperatures above the Ae<sub>1</sub> (1,330 °F). Below this temperature it decomposes into mixtures of ferrite and cementite (iron carbide). The end product or final structure is greatly influenced by the temperature at which the transformation occurs, and this, in turn, is influenced by the rate of cooling. Since the mechanical properties may be varied over a wide range, depending on the decomposition products of the parent austenite, a knowledge of how austenite decomposes and the factors influencing it is necessary for a clear understanding of the heat treatment of steel. The progressive transformation of austenite under conditions of equilibrium (extremely slow cooling) has been described. Practically, however, steel is not cooled under equilibrium conditions, and consequently the critical points on cooling always occur at lower temperatures than indicated in figure 4.

If samples of steel, say of eutectoid carbon content for the sake of simplicity, are cooled from above the Ae<sub>1</sub> at gradually increasing rates, the corresponding Ar transformation occurs at lower and lower temperatures (fig. 7). This transformation is distinguished from that occurring under extremely slow rates of cooling  $(Ar_1)$  by the designation Ar'. As the rate of cooling of this steel is increased, an additional transformation (termed the Ar" or

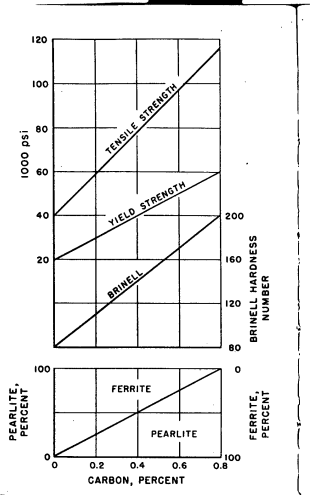
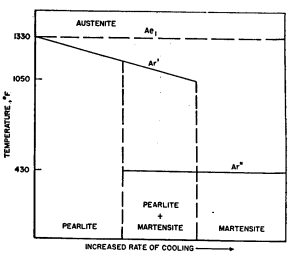
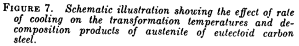


FIGURE 6. Relation of mechanical properties and structure to carbon content of slowly cooled carbon steels.





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 $M_{\bullet}$ ) appears at relatively low temperatures (about 430 °F). If the rate of cooling is still further increased, the Ar' transformation is suppressed entirely and only the Ar" transformation is evident. It should be noted that the temperature of the Ar" is not affected by the rate of cooling, whereas the temperature of the Ar' may be depressed to as low as about 1,050 °F in this particular steel.

The product of the Ar' transformation is fine pearlite. As the temperature of the Ar' is gradually lowered, the lamellar structure of the resulting pearlite becomes correspondingly finer and the steel becomes harder and stronger. The product of the Ar" transformation is martensite (fig. 9, A, and 15, A), which is the hardest and most brittle of the transformation products of austenite and is characterized by a typical acicular structure.

The phenomenon of the occurrence of both the Ar' and Ar" transformations is known as the split transformation. The resultant microstructures of steels cooled at such rates as to undergo a split transformation consist of varying amounts of fine pearlite and martensite (fig. 9, C). The actual amounts of these two constitutents are functions of the rates of cooling, the slower rates resulting in more pearlite and less martensite, and the faster rates resulting in more martensite and less pearlite.

#### 4.1. Isothermal Transformation

The course of transformation of austenite when the steel is quenched to and held at various constant elevated temperature levels (isothermal transformation) is conveniently shown by a diagram known as the S-curve (also termed the *TTT* diagram—for time, temperature, and transformation). Such a diagram for eutectoid carbon steel is shown in figure 8 and the discussion of this figure will be confined to steel of this particular composition.

#### a. To Pearlite

Austenite containing 0.80 percent of carbon, cooled quickly to and held at 1,300 °F, does not begin to decompose (transform) until after about 15 min, and does not completely decompose until after about 5 hr (fig. 8). Thus, at temperatures just below the Ae<sub>1</sub>, austenite is stable for a considerable length of time. The product of the decomposition of austenite at this temperature is coarse pearlite of relatively low hardness. If the austenite is quickly cooled to and held at a somewhat lower temperature, say 1,200 °F, decomposition begins in about 5 sec and is completed after about 30 sec, the resultant pearlite being finer and harder than that formed at 1,300 °F. At a temperature of about 1,050 °F, the austenite decomposes extremely rapidly, only about 1 sec elapsing before the transformation starts and 5

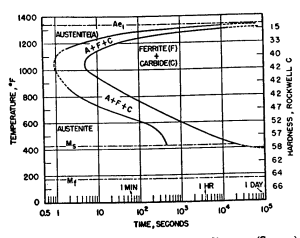


FIGURE 8. Isothermal transformation diagram (S-curve) for eutectoid carbon steel.

(Metals Handbook, 1948 edition, page 608.) The hardness of the structures formed at the various temperatures is given by the scale on the right.

sec before it is completed. The resultant pearlite is extremely fine and its hardness is relatively high. This region of the S-curve where decomposition of austenite to fine pearlite proceeds so rapidly is termed the "nose" of the curve.

#### b. To Bainite

If the austenite is cooled unchanged to temperatures below the nose of the S-curve  $(1,050 \, ^{\circ}\text{F})$ , the time for its decomposition begins to increase (fig. 8). The final product of decomposition now is not pearlite, but a new acicular constituent called bainite (fig. 15, E) possessing unusual toughness with hardness even greater than that of very fine pearlite.

Depending on the temperature, then, a certain finite interval of time is necessary before austenite starts to transform into either pearlite or bainite. Additional time is necessary before the transformations are completed.

#### c. To Martensite

If the austenite is cooled unchanged to a relatively low temperature (below about 430 °F for the eutectoid carbon steel under consideration), partial transformation takes place instantaneously; the product of transformation is martensite. Austenite transforms into mar-tensite over a temperature range and the amount that transforms is a function of the Only minute amounts will so temperature. transform at about 430 °F; practically all of the austenite will be transformed at about 175 °F. The beginning of this transformation range is termed the M<sub>s</sub> (martensitestart) and the end of the range is termed the  $M_t$  (martensite—finish). As long as the temperature is held constant within the  $M_s$ — $M_t$ range, that portion of the austenite that does not transform instantaneously to martensite

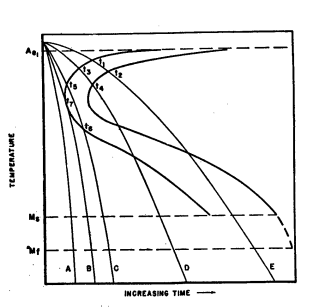
remains untransformed for a considerable length of time, eventually transforming to bainite.

In ordinary heat treatment of the plain carbon steels, austenite does not transform into bainite. Transformation of the austenite takes place either above or at the nose of the S-curve, forming pearlite, or in passing through the  $M_s$ — $M_t$  range, forming martensite or both. It is evident that in order for austenite to be transformed entirely into martensite, it must be cooled sufficiently rapidly so that the temperature of the steel is lowered past the nose of the S-curve in less time than is necessary for transformation to start at this temperature. If this is not accomplished, part of the steel transforms into pearlite at the high temperature (Ar'), and the remainder transforms into martensite at the low temperature (Ar'' or  $M_s$ —F, temperature range). This explains the phenomenon of the split transformation described previously.

#### 4.2. Continuous Cooling

Figure 9 represents a theoretical S-curve on which are superimposed five theoretical cooling curves. Curves A to E represent successively slower rates of cooling, as would be obtained, for instance, by cooling in iced brine, water, oil, air, and in the furnace, respectively.

The steel cooled according to curve E begins to transform at temperature  $t_1$  and completes transformation at  $t_2$ ; the final product is coarse pearlite with relatively low hardness. When cooled according to curve D, transformation begins at  $t_3$  and is completed at  $t_4$ ; the final product is fine pearlite and its hardness is greater than the steel cooled according to curve E. When cooled according to curve C, transformation begins at  $t_s$  and is only partially complete when temperature  $t_{6}$  is reached; the product of this partial transformation is very fine pearlite. The remainder of the austenite does not decompose until the M<sub>s</sub> temperature is reached, when it begins to transform to martensite, completing this transformation at the  $M_t$  temperature. The final structure is then a mixture of fine pearlite and martensite (typical of incompletely hardened steel-frequently termed "slack quenched" steel) with a higher hardness than was obtained with the steel cooled according to curve D. The rate of cooling represented by curve B is just sufficient to intersect the nose of the S-curve, consequently only a minute amount of the austenite decomposes into fine pearlite at temperature  $t_7$ ; the remainder of the austenite is unchanged until the martensite transformation range is reached. If the steel is cooled at a slightly faster rate, so that no transformation takes place at the nose of the S-curve, the steel is completely hardened. This particular rate is



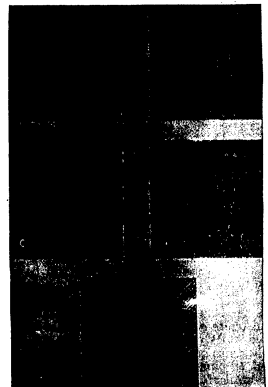


FIGURE 9. Schematic diagram illustrating the relation between the S-curve, continuous cooling curves, and resulting microstructures of eutectoid carbon steel.

Microstructures: A, martensite; B, martensite with a trace of very fine pearlite (dark); C, martensite and very fine pearlite; D, fine pearlite; E, coarse pearlite. All etched with nital.  $\times 600$ .

termed the critical cooling rate and is defined as the slowest rate at which the steel can be cooled and yet be completely hardened. Since this rate cannot be directly determined, the rate indicated by curve B, producing only a trace of fine pearlite (fig. 9, B), is frequently

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used as the critical cooling rate. The hardness of the resultant martensite is equivalent to the maximum that can be obtained. Samples cooled at a faster rate, such as that indicated by curve A, are also completely martensitic but no harder than the sample cooled according to the critical cooling rate.

It will be noted that the rate at which a steel cools through the temperature range in the vicinity of the nose of the S-curve is of critical importance. Somewhat slower rates of cooling above and below this temperature range can be tolerated and yet obtain a completely hardened steel, provided that the cooling through the temperature interval at the nose of the S-curve is sufficiently fast. In practice, however, steels are usually cooled rapidly from the quenching temperature to relatively low temperatures (about 500 °F) and then allowed to cool in air.

Although the above discussions of the decomposition of austenite have been limited to a steel of eutectoid composition, other steels behave in a similar manner, the temperatures and times of reactions being different. In hypoeutectoid steels, free ferrite plus pearlite are formed if transformation begins above the temperature range of the nose of the S-curve; the amount of free ferrite decreases as the temperature of transformation approaches the nose of the curve. In hypereutectoid steels, free cementite plus pearlite are formed if transformation occurs above the nose. time for the start of the transformation at the The nose increases as the carbon increases up to the eutectoid composition, and then decreases with further increase in carbon. That is, the nose is shifted to the right (with respect to the time axis, fig. 8) as the carbon is increased to 0.8 percent and to the left with further increase in carbon content.

The temperature of formation of bainite is not appreciably affected by carbon content, but the time for its formation increases with the carbon.

Both the  $M_s$  and  $M_t$  temperatures are markedly lowered by increasing carbon content, as is shown for  $M_s$  in figure 10. The  $M_t$ temperatures of the plain carbon steels have not been adequately determined; available information indicates that the  $M_t$  of high carbon steels is actually below room temperature. Slight amounts of austenite are frequently retained in quenched steels, especially in the higher carbon grades, even when cooled to room temperature.

## 5. Heat Treatment of Steels

All heat-treating operations consist of subjecting a metal to a definite time-temperature

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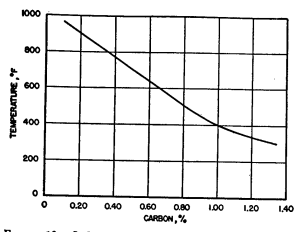


FIGURE 10. Influence of carbon on the start of the martensite  $(M_{\bullet})$  transformation of high-purity iron-carbon alloys.

(Digges, Trans. Am. Soc. Metals 28, 597 (1940).)

cycle, which may be divided into three parts: (1) Heating, (2) holding at temperature (soaking), and (3) cooling. Individual cases vary, but certain fundamental objectives may be stated.

The rate of heating is not particularly important unless a steel is in a highly stressed condition, such as is imparted by severe cold working or prior hardening. In such instances the rate of heating should be slow. Frequently this is impracticable, since furnaces may be at operating temperatures, but placing the cold steel in the hot furnace may cause distortion or even cracking. This danger can be minimized by the use of a preheating furnace maintained at a temperature below the A1. The steel, preheated for a sufficient period, then can be transferred to the furnace at operating temperature. This procedure is also advantageous when treating steels having considerable variations in section thickness or very low thermal conductivity.

The object of holding a steel at heat-treating temperature is to assure uniformity of temperature throughout its entire volume. Obviously, thin sections need not be soaked as long as thick sections, but if different thicknesses exist in the same piece, the period required to heat the thickest section uniformly governs the time at temperature. A rule frequently used is to soak  $\frac{1}{2}$  hr/in. of thickness.

The structure and properties of a steel depend upon its rate of cooling and this, in turn, is governed by such factors as mass, quenching media, etc. It must be realized that the thicker the section, the slower will be the rate of cooling regardless of the method of cooling used except in such operations as induction hardening to be discussed later.

#### 5.1. Annealing

Annealing is a process involving heating and cooling, usually applied to produce softening. The term also refers to treatments intended to alter mechanical or physical properties, produce a definite microstructure, or remove gases. The temperature of the operation and the rate of cooling depend upon the material being annealed and the purpose of the treatment.

#### a. Full Annealing

Full annealing is a softening process in which a steel is heated to a temperature above the transformation range  $(Ac_s)$  and, after being held for a sufficient time at this temperature, is cooled slowly to a temperature below the transformation range  $(Ar_1)$ . The steel is ordinarily allowed to cool slowly in the furnace, although it may be removed and cooled in some medium such as mica, lime, or ashes, that insures a slow rate of cooling.

Since the transformation temperatures are affected by the carbon content, it is apparent that the higher carbon steels can be fully annealed at lower temperatures than the lower carbon steels. In terms of the diagram shown in figure 4, steels must be heated to above the line GOSK. The temperature range normally used for full annealing is 25 to 50 deg F above this line (the upper critical), as shown in figure 11.

The microstructures of the hypoeutectoid steel: that result after full annealing are quite similar to those shown for the equilibrium conditions (fig. 5, B and C). Eutectoid and hypereutectoid steels frequently spheroidize partially or completely on full annealing (see later section and fig. 12, D).

#### b. Process Annealing

Process annealing, frequently termed stressrelief annealing, is usually applied to coldworked low carbon steels (up to about 0.25%of carbon) to soften the steel sufficiently to allow further cold-working. The steel is usually heated close to, but below, the Ac, temperature (fig. 11). If the steel is not to be further cold-worked, but relief of internal stresses is desired, a lower range of temperature will suffice (about 1,000 °F). Rate of cooling is immaterial.

This type of anneal will cause recrystallization and softening of the cold-worked ferrite grains, but usually will not affect the relatively small amounts of cold-worked pearlite. Typical structures of cold-worked, process-annealed, and fully annealed low-carbon steel are shown in figure 12, A, B, and C, respectively.

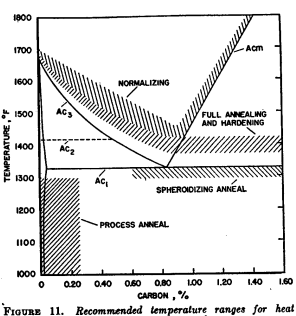


FIGURE 11. Recommended temperature ranges for heat treating plain carbon steels.

#### c. Spheroidizing

Spheroidizing is a process of heating and cooling steel that produces a rounded or globular form of carbide in a matrix of ferrite. It is usually accomplished by prolonged heating at temperatures just below the Ac<sub>1</sub> (fig. 11), but may be facilitated by alternately heating to temperatures just above the Ac<sub>1</sub> and cooling to just below the Ar<sub>1</sub>. The final step, however, should consist of holding at a temperature just below the critical (Ar<sub>1</sub>). The rate of cooling is immaterial after slowly cooling to about 1,000 °F.

The rate of spheroidization is affected by the initial structure. The finer the pearlite, the more readily spheroidization is accomplished. A martensitic structure is very amenable to spheroidization.

This treatment is usually applied to the high carbon steels (0.60% of carbon and higher). The purpose of the treatment is to improve machinability and it is also used to condition high-carbon steel for cold-drawing into wire. A typical microstructure of spheroidized high carbon steel is shown in figure 12, D.

#### 5.2. Normalizing

Normalizing is a process in which a steel is heated to a temperature above the  $A_{c_3}$  or the  $A_{cm}$  (fig. 11) and then cooled in still air. The purpose of the treatment is to obliterate the effects of any previous heat treatment (including the coarse-grained structure sometimes

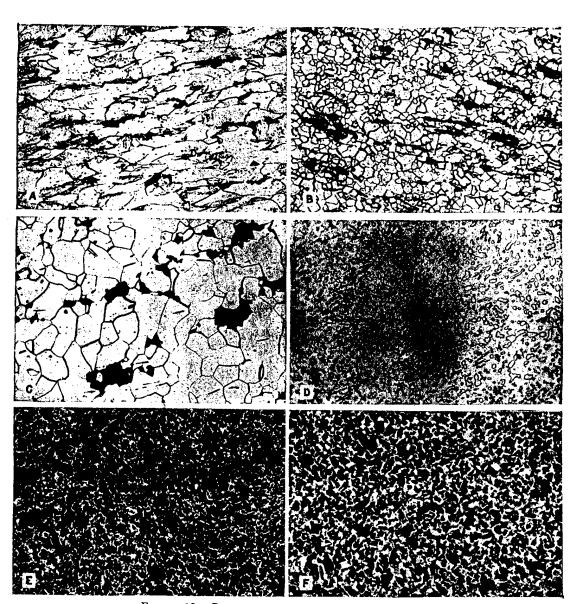


FIGURE 12. Representative microstructures of carbon steels.

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FIGURE 12. Kepresentative microstructures of carbon steels. A, Low carbon steel (0.1% C) as cold-worked. Both ferrite (light) and pearlite (dark) are severely deformed. ×100. B, Bame as (A) after process annealing at 1,200° F. The ferrite is recrystallized (grains are equi-ared), but pearlite is not affected by this treatment. ×100. C, Same as (A) after full annealing at 1,600° F. The ferrite is recrystallized (grains are equi-ared), but pearlite is not affected by this treatment. ×100. D, High carbon steel (1.1% C) as spheroidized. All traces of cold working are eliminated, and the ferrite grains are larger than in (B). ×100. E, Hypoeutectoid steel (0.5% C) as normalized at 1,600° F in ¼-in. round (center area). Because of the rapid rate of air cooling such a small section, the pearlite is quite fine and relatively little free ferrite is formed. ×100. F, Same steel as (E) but normalized in 2¼-in. round (center area). The slower rate of cooling due to the larger section results in coarser pearlite and more free ferrite. ×100. All etched in picral.

resulting from high forging temperatures) or cold-working and to insure a homogeneous austenite on reheating for hardening or full annealing. The resultant structures are pearl-ite or pearlite with excess ferrite or cementite, depending upon the composition of the steel. They are different from the structures resulting after annealing in that, for steels of the same carbon content in the hypo- or hypereutectoid ranges, there is less excess ferrite or

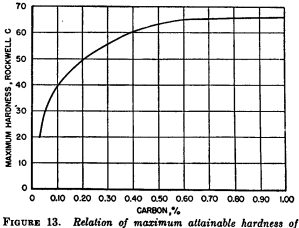
cementite and the pearlite is finer. These are the results of the more rapid cooling.

Since the type of structure, and, therefore, the mechanical properties, are affected by the rate of cooling, considerable variations may occur in normalized steels because of differences in section thickness of the shapes being normalized. The effect of section thickness on the structure of a normalized 0.5-percentcarbon steel is shown in figure 12, E and F.

#### 5.3. Hardening

Steels can be hardened by the simple expedient of heating to above the Ac, transformation. holding long enough to insure the attainment of uniform temperature and solution of carbon in the austenite, and then cooling rapidly (quenching). Complete hardening depends on cooling so rapidly (fig. 9, A and B) that the austenite, which otherwise would decompose on cooling through the Ar<sub>1</sub>, is maintained to relatively low temperatures. When this is accomplished, the austenite transforms to martensite on cooling through the M<sub>s</sub>-M<sub>f</sub> range. Rapid cooling is necessary only to the extent of lowering the temperature of the steel to well below the nose of the S-curve. Once this has been accomplished, slow cooling from then on, either in oil or in air, is beneficial in avoiding distortion and cracking. Special treatments, such as time quenching and martempering, are designed to bring about these conditions. As martensite is quite brittle, steel is rarely used in the as-quenched condition, that is, without tempering.

The maximum hardness that can be obtained in completely hardened low-alloy and plain carbon structural steels depends primarily on the carbon content. The relationship of maximum hardness to carbon content is shown in figure 13.



quenched sleels to carbon content. (Burns, Moore, and Archer, Trans. Am. Soc. Metals 26, 14 (1938)).

#### a. Effect of Mass

Previous discussion of the formation of martensite has neglected the influence of mass. It must be realized that even with a sample of relatively small dimensions, the rate of abstraction of heat is not uniform. Heat is always abstracted from the surface layers at a faster rate than from the interior. In a given cooling medium the cooling rate of both the surface and interior decreases as the dimensions of a

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sample increase and the possibility of exceeding the critical cooling rate (fig. 9, B) becomes less. To overcome this, the steel may be quenched in a medium having a very high rate of heat abstraction, such as iced brine, but, even so, many steels have a physical restriction on the maximum size amenable to complete hardening regardless of the quenching medium.

The marked effect that mass has upon the hardness of quenched steel may be illustrated by measuring the hardness distribution of different size rounds of the same steel quenched in the same medium. Curves showing the distribution of hardness in a series of round bars of different sizes of 0.5-percent-carbon steel are shown in figure 14. The quenching medium used was water; the quenching temperature was 1,530 °F. The rate of cooling decreased as the diameters of the bars increased. Only the  $\frac{1}{2}$ -in. round hardened completely through the cross section, whereas with the 4-in. round the critical cooling rate was not attained even at the surface.

#### 5.4. Tempering

Tempering (sometimes called drawing) is the process of reheating hardened (martensitic) or normalized steels to some temperature below the lower critical ( $Ac_1$ ). The rate of cooling

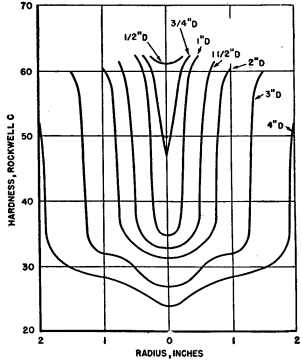


FIGURE 14. Variation in hardness in different size rounds of 0.5-percent-carbon steels as quenched from 1,530° F in water.

(Jominy, Hardenability of alloy steels, Am. Soc. Metals, p. 75, 1939.)

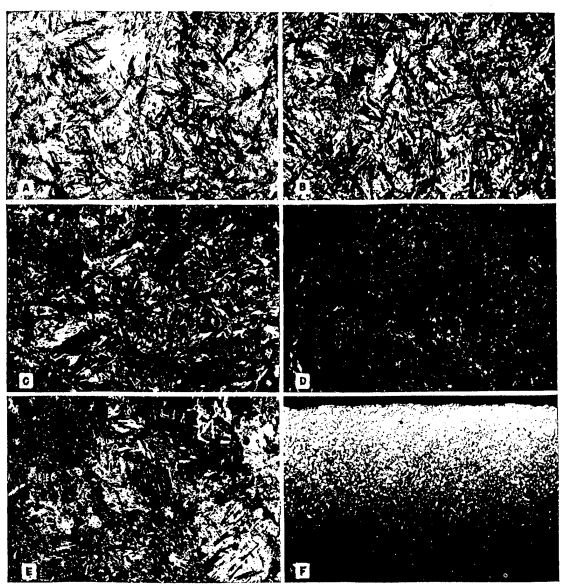


FIGURE 15. Microstructures and corresponding hardness of heat-treated high carbon steel.

A, B

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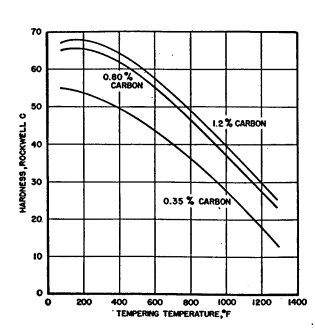
A, As quenched in brine; martensite, Rockwell C hardness=50, X500. B, Same as A after tempering at 400° F; tempered martensite, Rockwell C hardness=61. X500. C, Same as (A) after tempering at 800° F; tempered martensite, Rockwell C hardness=45. X500. D, Same as (A) after tempering at 1,200° F; tempered martensite, Rockwell C hardness=25. X500. E, Quenched in lead at 650° F; bainite, Rockwell C hardness=49. X500. F, Heat-treated steel showing decarburized surface layer (light area) that did not respond to hardening. X100. All etched in 1% nital.

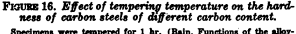
is immaterial except for some steels that are susceptible to temper brittleness.<sup>2</sup>

As the tempering temperature is increased. the martensite (fig. 15, A) of hardened steel passes through stages of tempered martensite (fig. 15, B and C) and is gradually changed

into a structure consisting of spheroids of cementite in a matrix of ferrite, formerly termed sorbite (fig. 15, D). These changes are accompanied by a decreasing hardness and increasing toughness. The tempering temperature depends upon the desired properties and the purpose for which the steel is to be used. If considerable hardness is necessary, the tempering temperature should be low; if consid-erable toughness is required, the tempering temperature should be high. The effect of tempering on the hardness of fully hardened carbon steels is shown in figure 16.

<sup>&</sup>lt;sup>2</sup> Temper brittleness is manifested as a loss of toughness (ob-served only by impact tests of notched bars) after slow cooling from tempering temperatures of 1,100 °F or higher, or after tem-pering in the temperature range between approximately 850 ° and 1,100 °F. It is most pronounced in alloy steels that contain manganese or chromium and usually can be prevented by rapid quenching from the tempering temperature. The presence of molyb-denum is beneficial in counteracting the tendency toward temper brittleness. brittleness





Specimens were tempered for 1 hr. (Bain, Functions of the alloying elements in stel, Am. Soc. Metals, p. 38, 1939.)

Proper tempering of a hardened steel requires a certain amount of time. At any selected tempering temperature, the hardness drops rapidly at first, gradually decreasing more slowly as the time is prolonged. The effect of time at different tempering temperatures upon the resultant hardness of a eutectoid carbon steel is shown in figure 17. Short tempering periods are generally undesirable and should be avoided. Good practice requires at least  $\frac{1}{2}$  hr (or, preferably, 1 to 2 hr) at tempering temperature for any hardened steel.

The necessity for tempering a steel promptly after hardening cannot be overemphasized. If fully hardened steel is allowed to cool to room temperature during hardening there is danger that the steel may crack. Carbon steels and most of the low alloy steels should be tempered as soon as they are cool enough to feel comfortable (100 to 140 °F) to the bare hands. Steels should not be tempered before they cool to this temperature range because in some steels the M<sub>t</sub> temperature is quite low and untransformed austenite may be present. Part or all of this residual austenite will transform to martensite on cooling from the tempering temperature so that the final structure will consist of both tempered and untempered martensite. The brittle untempered martensite, together with the internal stresses caused by its formation, can easily cause failure of the heat-treated part. When it is possible that such a condition exists, a second tempering treatment (double tempering) should be given to temper the fresh martensite formed on cooling after the initial tempering treatment.

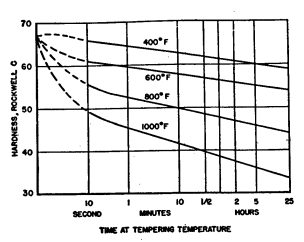


FIGURE 17. Effect of time at tempering temperature on the hardness of 0.8-percent-carbon steel. (Logarithmic time scale.)

(Bain, Functions of the alloying elements in steel, Am. Soc. Metals, p. 233, 1939.)

If structural steels are to be used in the normalized condition, the normalizing operation is frequently followed by heating to a temperature of about 1,200 to 1,300 °F. The purpose of this treatment, which is also designated as tempering, or stress-relief annealing, is to relieve internal stresses resulting on cooling from the normalizing temperature and to improve the ductility.

## 5.5. Case Hardening

Case hardening is a process of hardening a ferrous alloy so that the surface layer or case is made substantially harder than the interior or core. The chemical composition of the surface layer is altered during the treatment by the addition of carbon, nitrogen, or both. The most frequently used case-hardening processes are carburizing, cyaniding, carbonitriding, and nitriding.

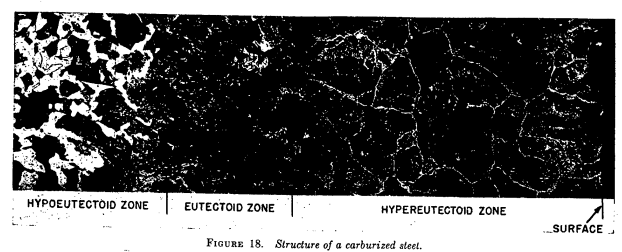
#### a. Carburizing

Carburizing is a process that introduces carbon into a solid ferrous alloy by heating the metal in contact with a carbonaceous material to a temperature above the  $Ac_3$  of the steel and holding at that temperature. The depth of penetration of carbon is dependent on temperature, time at temperature, and the composition of the carburizing agent. As a rough indication, a carburized depth of about 0.030 to 0.050 in. can be obtained in about 4 hr at 1,700 °F, depending upon the type of carburizing agent, which may be a solid, liquid, or gas.

Since the primary object of carburizing is to secure a hard case and a relatively soft, tough core, only low-carbon steels (up to a maximum of about 0.25% of carbon), either with or without alloying elements (nickel, chromium, man-

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The specimen was cooled slowly from the carburizing temperature. Etched with nital. X75.

ganese, molybdenum), are normally used. After carburizing, the steel will have a highcarbon case graduating into the low-carbon core (fig. 18).

A variety of heat treatments may be used subsequent to carburizing, but all of them involve quenching the steel to harden the carburized surface layer. The most simple treatment consists of quenching the steel directly from the carburizing temperature; this treatment hardens both the case and core (insofar as the core is capable of being hardened). Another simple treatment, and perhaps the one most frequently used, consists of slowly cooling from the carburizing temperature, reheating to above the Ac<sub>3</sub> of the case (about 1,425 °F). and quenching; this treatment hardens the case only. A more complex treatment is to double quench—first from above the Ac<sub>3</sub> of the core (about 1,650 °F for low-carbon steel) and then from above the Ac<sub>3</sub> of the case (about 1,425 °F); this treatment refines the core and hardens the case. The plain carbon steels are almost always quenched in water or brine; the alloy steels are usually quenched in oil. Although tempering following hardening of carburized steel is sometimes omitted, a low-temperature tempering treatment at about 300 °F is good practice.

It is sometimes desirable to carburize only certain parts of the surface. This can be accomplished by covering the surface to be protected against carburizing with some material that prevents the passage of the carburizing agent. The most widely used method is copper plating of the surfaces to be protected. Several proprietary solutions or pastes, which are quite effective in preventing carburization, are also available.

The commercial compounds commonly used for pack (solid) carburizing contain mixtures of carbonate (usually barium carbonate), coke (diluent), and hardwood charcoal, with oil, tar,

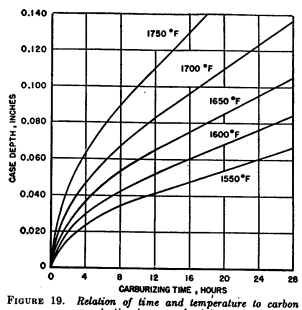
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or molasses as a binder. Mixtures of charred leather, bone, and charcoal are also used.

The carburizing action of these compounds is diminished during use and it is necessary to add new material before the compound is reused. Addition of one part of unused to three to five parts of used compound is common practice. The parts to be carburized are packed in boxes (or other suitable containers) made of heat-resistant alloys, although rolled or cast steel may be used where long life of the box is not important. The lid of the box should be sealed with fire clay or some other refractory to help prevent escape of the carburizing gas generated at the carburizing temperature. The depth and uniformity of case is affected by the method of packing and design of the container.

Liquid carburizing consists of case hardening steel or iron in molten salt baths that contain mixtures principally of cyanides (poisonous), chlorides, and carbonates. The case produced by this method contains both carbon and nitrogen, but principally the former. The temperatures used range from about 1,550 to 1,650 °F or higher, depending upon the compositions of the bath and the desired depth of case. At 1,650 °F a case depth of about 0.010 to 0.015 in. can be obtained in 1 hr; about 0.020 to 0.030 in. can be obtained in 4 hr. Considerably deeper cases can be obtained at higher temperatures with longer periods of time. After carburizing, the parts must be quenched just as in solid carburizing, but it is usual to do this directly from the molten bath.

In all present-day commercial gas carburizing, two or more hydrocarbons are used in combination for supplying the carbon to the steel. The hydrocarbons used are methane, ethane, propane, and oil vapors. The steel parts are placed in sealed containers through which the carburizing gases are circulated; the temperatures used are in the neighborhood of



penetration in gas carburizing. (Metals Handbook, 1939 edition, p. 1041.)

1,700 °F. Average expectation for depth of case in gas-carburized steel is illustrated in figure 19. After carburizing, the parts must be quench hardened.

#### b. Cyaniding

A hard, superficial case can be obtained rapidly on low-carbon steels by cyaniding. This process involves the introduction of both carbon and nitrogen into the surface layers of the steel. Steels to be cyanided normally are heated in a molten bath of cyanide-carbonatechloride salts (usually containing 30 to 95% ot sodium cyanide) and then quenched in brine, water, or mineral oil; the temperature of operation is generally within the range of 1,550 to 1,600 °F. The depth of case is a function of time, temperature, and composition of the cyanide bath; the time of immersion is guite short as compared with carburizing, usually varying from about 15 min to 2 hr. The maximum case depth is rarely more than about 0.020 in. and the average depth is considerably less

Steels can be cyanided also by heating to the proper temperature and dipping in a powdered cyanide mixture or sprinkling the powder on the steel, followed by quenching. The case thus formed is extremely thin.

Cyaniding salts are violent poisons if allowed to come in contact with scratches or wounds; they are fatally poisonous if taken internally. Fatally poisonous fumes are evolved when cyanides are brought into contact with acids. Cyaniding baths should be equipped with a hood for venting the gases evolved during heating and the work room should be well ventilated. Molten cyanide should never be permitted to come in contact with sodium or or potassium nitrates commonly used for baths for tempering as the mixtures are explosive. Furthermore, care is necessary in preparing a salt bath and the work pieces should be completely dry before placing in the molten bath. The advice of salt manufacturers should be obtained and followed in the operation and maintenance of salt baths.

#### c. Carbonitriding

Carbonitriding, also termed gas cyaniding, dry cyaniding, and nitrocarburizing, is a process for case hardening a steel part in a gascarburizing atmosphere that contains ammonia in controlled percentages. Carbonitriding is used mainly as a low-cost substitute for cyaniding and, as in cyaniding, both carbon and nitrogen are added to the steel. The process is carried on above the Ac1 temperature of the steel, and is practical up to 1,700 °F. Quenching in oil is sufficiently fast to attain maximum surface hardness; this moderate rate of cooling tends to minimize distortion. The depth to which carbon and nitrogen penetrate varies with temperature and time. The penetration of carbon is approximately the same as that obtained in gas carburizing (fig. 19).

#### d. Nitriding

The nitriding process consists in subjecting machined and heat-treated steel, free from surface decarburization, to the action of a nitrogenous medium, usually ammonia gas, at a temperature of about 950 to 1,050 °F, whereby a very hard surface is obtained. The surface-hardening effect is due to the absorption of nitrogen and subsequent heat treatment of the steel is unnecessary. The time required is relatively long, normally being 1 or 2 days. The case, even after 2 days of nitriding, is generally less than 0.020 in. and the highest hardness exists in the surface layers to a depth of only a few thousandths of an inch.

Liquid nitriding (nitriding in a molten salt bath, generally composed of a mixture of sodium and potassium salts) employs the same temperature range as gas nitriding.

Special low-alloy steels have been developed for nitriding. These steels contain elements that readily combine with nitrogen to form nitrides, the most favorable being aluminum, chromium, and vanadium. Molybdenum and nickel are used in these steels to add strength and toughness. The carbon content usually is about 0.20 to 0.50 percent, although in some steels, where high core hardness is essential, it may be as high as 1.3 percent. Stainless steels also can be nitrided.

Because nitriding is carried out at a relatively low temperature, it is advantageous to

use quenched and tempered steel as the base material. This gives a strong, tough core with an intensely hard wear-resisting case—much harder, indeed, than can be obtained by quench hardening either carburized or cyanided steel.

Although warpage is not a problem during nitriding, steels increase slightly in size during this treatment. Allowance can be made for this growth in the finished article. Protection against nitriding can be effected by tin, copper, or bronze plating, or by the application of certain paints.

#### 5.6. Surface Hardening

It is frequently desirable to harden only the surface of steels without altering the chemical composition of the surface layers. If a steel contains sufficient carbon to respond to hardening, it is possible to harden the surface layers only by very rapid heating for a short period, thus conditioning the surface for hardening by quenching.

#### a. Induction Hardening

In induction hardening, a high-frequency current is passed through a coil surrounding the steel, the surface layers of which are heated by electro-magnetic induction. The depth to which the heated zone extends depends on the frequency of the current (the lower frequencies giving the greater depths) and on the duration of the heating cycle. The time required to heat the surface layers to above the  $Ac_3$  is surprisingly brief, frequently being a matter of only a few seconds. Selective heating (and therefore hardening) is accomplished by suitable design of the coils or inductor blocks. At the end of the heating cycle, the steel is usually quenched by water jets passing through the inductor coils. Precise methods for controlling the operation, that is, rate of energy input, duration of heating, and rate of cooling, are necessary. These features are incorporated in induction hardening equipment, which is usually entirely automatic in operation. Figure 20 shows the hardened layer in induction hardened steel.

#### b. Flame Hardening

Flame hardening is a process of heating the surface layers of steel above the transformation temperature by means of a high-temperature flame and then quenching. In this process the gas flames impinge directly on the steel surface to be hardened. The rate of heating is very rapid, although not so fast as with induction heating. Plain carbon steels are usually quenched by a water spray, whereas the rate of cooling of alloy steels may be varied from a rapid water quench to a slow air cool depending on the composition.

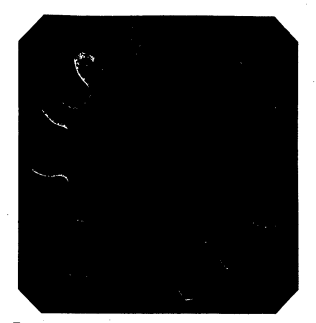


FIGURE 20. Macrograph of induction hardened gear teeth. Original size, 4-in. diameter (Osborn).

Any type of hardenable steel can be flame hardened. For best results, the carbon content should be at least 0.35 percent, the usual range being 0.40 to 0.50 percent.

#### 5.7. Special Treatments

#### a. Austempering

Austempering is a trade name for a patented heat-treating process. Essentially, it consists of heating steel to above the  $Ac_3$  transformation temperature and then quenching into a hot bath held at a temperature below that at which fine pearlite would form (the nose of the S-curve, fig. 8), but above the M<sub>s</sub> temperature (fig. 10). The product of isothermal decomposition of austenite in this temperature region is bainite. This constituent combines relatively high toughness and hardness. A typical microstructure of austempered steel is shown in figure 15, E.

The austempering process has certain limitations that make it impracticable for use with many steels. In order to assure a uniform structure (and hence uniform properties), it is essential that the entire cross section of the steel be cooled rapidly enough so that even the center escapes transformation at the nose of the S-curve. In carbon steels the time required to start transformation at the nose of the S-curve is extremely short, so that only relatively small sections (about  $\frac{3}{8}$ -in. maximum thickness) can be successfully hot quenched in austempering baths. The time required for transformation of the austenite of alloy steels

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to fine pearlite is usually longer, and hence larger sections can be successfully austempered (about 1 in. maximum). However, the time required for transformation to bainite frequently becomes inordinately long with many alloy steels and the process of austempering, therefore, is generally impracticable for these steels.

#### b. Martempering

Martempering consists of heating a steel to above its  $Ac_s$  transformation and then quenching into a bath held at a temperature approximately equal to that of its  $M_s$ . The steel is maintained in the hot bath until its temperature is essentially uniform and then is cooled in air.

Severe internal stresses develop in steel during hardening. Steel contracts during cooling but undergoes a marked expansion when the austenite transforms to martensite. Since a quenched steel must cool from the surface inward, various portions transform at different times. The steel is thus subjected to a variety of differential expansions and contractions, resulting in considerable internal stress. By equalizing the temperature throughout the section before transformation takes place, and then cooling slowly through the martensite  $(M_s - M_t)$  range, the internal stresses are considerably reduced. This is the prime objective of martempering.

Modified martempering differs from standard martempering only in that the bath temperature is lower, ranging from just below the M. point to about 200 °F. With the faster cooling at these lower bath temperatures, steels of lower hardenability can be hardened to sufficient depth. However, greater distortion of sensitive parts is likely with the modified process.

#### c. Cold Treatment

The M<sub>t</sub> temperature of many alloy steels is so low that complete transformation of austenite to martensite does not occur on quenching to room temperature or on cooling after tempering. This retained austenite may be partially or completely transformed by cooling below atmospheric temperatures and such treatment is called "cold treatment." The beneficial effects of cold treatment have not been fully explored. It is known that the retained austenite of highly alloyed steels is frequently difficult to transform. Cooling these steels to low temperatures (to the temperature of solid  $CO_2$  or even lower) immediately after the quench is sometimes effective in transforming the retained austenite, but with the concomitant danger of cracking. When the cold treatment is applied after tempering, the retained austenite is considerably more resistant to

transformation. If cold treatment is used, the steel should always be tempered afterwards.

Repeated alternate heating to a temperature slightly below that used in tempering and cooling to a subzero temperature in a refrigerated iced brine, carbon dioxide, liquid air, or liquid nitrogen is commonly used for transforming the retained austenite (dimensional stabilization) of steel gages, especially those of the ballbearing type composition (AISI 52100).

#### d. Ausforming

In this process, medium-carbon alloy steels are first austenitized and then cooled rapidly to the temperature range above M, in the "bay" of the S-curve, between the pearlite and bainite transformation bands. While held at a temperature within this bay the steels are plastically deformed and subsequently transformed at lower temperatures to martensite or bainite. The steels are then tempered. This "ausforming" or "hot-working" technique has been employed to provide higher yield and tensile strength values than those obtainable for these steels by the normal quench and temper treatments.

#### 6. Hardenability

Hardenability is the property that determines the depth and distribution of hardness induced in steel by quenching. It is increased by increasing carbon and by the addition of all the common alloying elements (except cobalt), provided that these elements are completely dissolved in the austenite at quenching temperatures. The elements most frequently used for this purpose are manganese, chromium, and molybdenum. Hardenability is also enhanced by increased grain size<sup>3</sup> and homogeneity of the austenite. However, a coarse-grained austenite increases the tendency of a steel to

<sup>3</sup> The grain size that influences hardenability is that grain size of the austenite that exists at the quenching temperature. It is usually measured under the microscope in terms of the number of grains per square inch at a magnification of  $\times 100$ . The common range of grain size numbers is as follows (note that the larger the number, the finer is the grain size; i.e., the more grains there are per square inch):

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		-	
Grain size no.	Grains/in <sup>2</sup> at ×100	Grain size	Grains/in <sup>3</sup> at ×100
0	0.50	7.5	90.5
0.5	0.707	8.0	128.
1.0			
		8.5	181.
1.5	1.41	9.0	256.
2.0	2.0	9.5	362.
2.5	2.83		
8.0			
	4.0	10.5	724.
8.5	5.66	11.0	
4.0	8.0		
4.5			
	11.8	12.0	2050.
5.0	16.0	12.5	2900.
5.5	22.6	13.0	
6.0	32.0		
		13.5	5800.
6.5	45.3	14.0	
7.0	610		

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distort and crack during heat treatment. Coarse-grained steels also are less tough than fine-grained steels.

A clear distinction must be drawn between the maximum hardness obtainable in a steel and its hardenability. In straight carbon and low-alloy steels, the maximum hardness is a function of carbon content only (fig. 13), whereas hardenability is concerned primarily with the depth of hardening.

Numerous methods have been proposed and used for determining the hardenability of steel. The selection of a method depends largely upon the information desired and the range in hardenability of the steels.

For a fundamental study, hardenability is often measured in terms of the critical cooling rate (fig. 9, B), but the standard end-quench (Jominy) test is widely used commercially. In the end-quench test a bar of steel 1 in. in diameter by 4 in. long, previously normalized and machined to remove the decarburized surface, is heated to the hardening temperature for 30 min. It is then quickly transferred to a fixture that holds the bar in a vertical position and a jet of water under controlled conditions immediately is directed against the bottom end only. The end of the bar is thus cooled very quickly while cross sections remote from the end are cooled more slowly, the rate of cooling being dependent upon the distance from the quenched end. After cooling is completed, two diametrically opposite flats about 1/4 in. wide are ground the length of the bar and Rockwell hardness measurements are made at intervals of  $\frac{1}{16}$  in. on one or both of the flat surfaces so prepared. The relationship of hardness to distance from the quenched end is an indication of the hardenability (fig. 21)

The conditions of the end-quench test have been standardized and the rates of cooling at various distances from the quenched end have been determined and are shown at the top of figure 21. It is thus possible to correlate hardness with cooling rate, which is a function of the distance from the water-cooled end. The cooling rates at various positions in the crosssections of different size rounds, when quenched in still oil and in still water also have been determined. Correlations can be made, therefore, between the hardness obtained at various positions on the end-quench bar and positions of equivalent hardness of round bars quenched in still oil or water (assuming that points of equal hardness have the same cooling rate). Figure 22 shows this relationship between the distances on the end-quenched bar and various locations for equivalent hardnesses in round bars quenched in still oil; figure 23 shows corresponding relations for bars quenched in still water.

If the hardenability (end-quench) curve of a steel is known, it is possible to ascertain

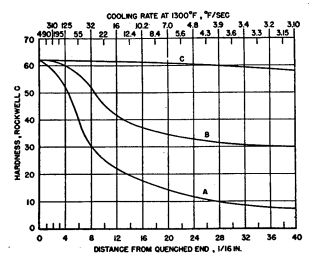


FIGURE 21. Hardenability curves for different steels with the same carbon content.

A, Shallow hardening; B, intermediate hardening; C, deep hardening. The cooling rates shown in this figure are according to the results of Boegehold and Weinman, N.D.R.C. Report OSRD No. 3743, p. 42 (June 1, 1944).

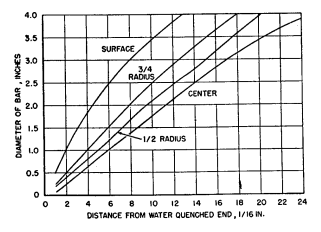
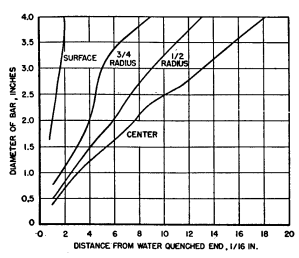
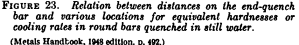


FIGURE 22. Relation between distances on the end-quench bar and various locations for equivalent hardness or cooling rates in round bars quenched in still oil. (Metals Handbook, 1948 edition, p. 492.)

from this curve and the curves given in figures 22 and 23 whether or not a selected size round made from the steel will have a desired hardness at its surface, <sup>3</sup>/<sub>4</sub> radius, <sup>1</sup>/<sub>2</sub> radius, or center, when quenched either in still oil or water. Conversely, the hardness values as determined at various positions in rounds after quenching in oil or water can be used to advantage in approximating the degree of hardenability of a steel suitable for replacement of similar parts.

The standard end-quench test is most useful for steels of moderate hardenability, that is, the low-alloy steels. For shallow hardening steels, such as plain carbon, a modification of this test (the L-type bar) is used.





#### 7. Heat Treatment of Cast Irons

Cast irons are of three general types, gray, white, and nodular. The terms gray and white are descriptive of the characteristic appearances of the fractures and nodular is descriptive of the shape of the graphite particles.

In gray iron a large proportion of the carbon is present as graphitic or uncombined carbon in the form of flakes. The small proportion of carbon that is present as combined carbon or cementite is in the form of pearlite just as in a steel. Depending on the amount of combined carbon, the matrix of cast iron may be similar to a hypoeutectoid steel with varying amounts of free ferrite and graphite flakes dispersed throughout (fig. 24, A, B, and C). If the amount of combined carbon is about 0.8 percent, the matrix will be entirely pearlitic; if the amount is greater, the matrix will be similar to a hypereutectoid steel.

In white iron most, if not all, of the carbon is present as cementite (fig. 24, D). The dividing line between white and gray cast iron is not clearly defined and iron possessing characteristics of both are called mottled cast iron.

Whether a cast iron will be gray or white depends on two factors—composition and rate of cooling. Certain elements tend to promote the formation of graphitic carbon; the most important of these are silicon, nickel, and sulfur. Others tend to promote the formation of combined carbon; the most important of these are chromium, molybdenum, vanadium, and manganese. Rapid solidification and cooling, such as result from thin sections cast in sand or larger sections cast against chills, promote the formation of white iron. Iron so produced is frequently termed chilled iron. Because of variations in the rate of solidification and cooling, castings may be white at the surface layers, mottled immediately below, and gray in the interior.

In nodular cast iron, also known as "ductile iron," "nodular graphite iron," "spherulitic iron," and "spheroidal graphite iron," the graphite is present in the form of nodules or spheroids (fig. 24, E and F) bestowing considerable ductility to the cast iron. These nodules form during solidification due to the treatment of the molten iron, just prior to casting, with a few hundredths percent usually of either magnesium or cerium; the sulfur content must be below about 0.015 percent for the treatment to be effective with these elements. Nodular iron and gray iron have the same total carbon content. The mechanical properties of nodular iron can be varied by alloying.

Cast irons may be given a variety of heat treatments, depending on the compositions and desired properties. The principles of the heat treatments applied to cast iron are similar to those already discussed for steels.

#### 7.1. Relieving Residual Stresses (Aging)

The heat treatment that relieves residual stresses is commonly called "aging," "normalizing," or "mild annealing;" the term "stress relieving" is more accurately descriptive. This treatment can be accomplished by heating the cast iron to between 800 and 1,100 °F., holding at temperature from 30 min. to 5 hr, the time depending on the size and temperature, and cooling slowly in the furnace. Such treatment will cause only a slight decrease in hardness, very little decomposition of cementite, and only a slight change in the strength of the metal.

#### 7.2. Annealing

Sometimes, it is desirable to soften castings in order to facilitate machining. The temperature range most commonly used for this purpose is 1,400 ° to 1,500 °F, although temperatures as low as 1,200 ° to 1,250 °F have been used satisfactorily. Highly alloyed irons are sometimes annealed at temperatures as high as 1,800 °F. In all annealing, care should be taken to prevent oxidation of the casting and cooling should be slow.

In general, softening for machinability is accompanied by a decrease in strength and in the amount of combined carbon, and by an increase in graphite content. Irons that contain carbide-forming elements (such as chromium, molybdenum, vanadium, and manganese) are more resistant to softening than the ordinary gray iron and, therefore, must be annealed at considerably higher temperatures. Completely annealed ordinary gray cast iron has a Brinell

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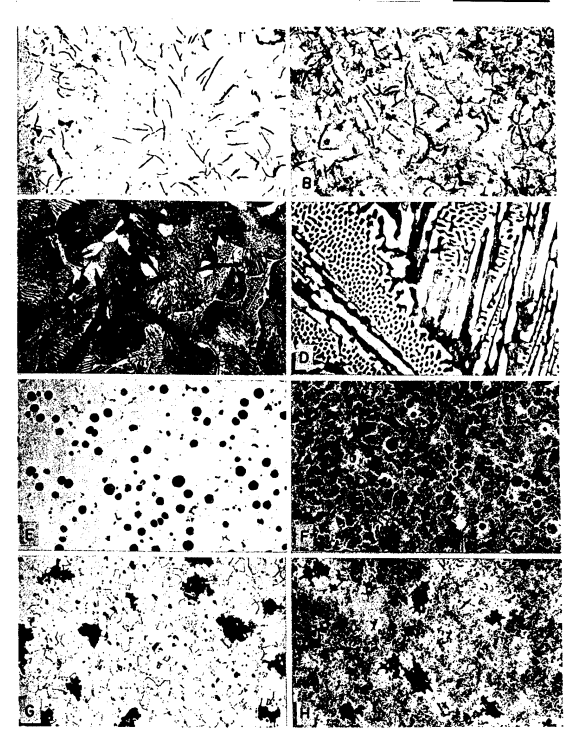


FIGURE 24. Microstructure of cast irons.

A. Gray iron. The dark flakes are graphite. Unetched. ×100.
B. Same as (A). Etched with picral. ×100.
C. Same as (B). The matrix consists of pearlite with a small amount of free ferrite (white) and is similar to a hypoeutectoid steel. The appearance of the graphite flakes is not affected by the etchant. ×500.
D. White iron. The white needles are cementite; the dark areas are pearlite; the dark spheroidal areas are the eutectic (Ledeburite). Etched with picral. ×500.
E. Nodular iron. The dark nodules are graphite; the white matrix is ferrite. Etched with picral. ×100.
F. Pearlitic nodular iron. Nodular graphite (dark) in a pearlitic matrix. Etched with picral. ×100.
H. Pearlitic malleable iron. Nodular graphite (dark) in a pearlite matrix. Etched with picral. ×100.

hardness of 120 to 130, and completely annealed alloy irons may have a Brinell hardness of 130 to 180, depending on the composition.

The matrix of nodular iron can be made completely ferritic by heating at 1,300 °F for 1 to 5 hr, depending on composition, size of casting, and amount of pearlite in the as-cast structure; cooling from the annealing temperature can be at any convenient rate. If massive or primary carbides are present in the as-cast structure, they may be converted to spheroidal graphite by heating at 1,600 to 1,700 °F for 1 to 5 hr. Slowly cooling from this temperature range to about 1,275 °F and holding from 3 to 5 hr will convert the matrix to ferrite. Nodular iron with a ferritic matrix has the desirable combination of maximum machinability, toughness, and ductility. The hardness of ferritic nodular iron varies with the alloy content from a low of about 130 Brinell to a high of about 210 Brinell.

#### a. Malleabilizing

Malleabilizing (also termed graphitizing) is a process of annealing white cast iron in such a way that the combined carbon is wholly or partly transformed to graphite in the nodular form (fig. 24, G and H) instead of the flaky form that exists in gray cast iron. The nodular form of carbon is called temper or free carbon. In some instances, part of the carbon is lost in the malleabilizing process.

White iron that is to be malleabilized should contain all its carbon in the combined form. Any carbon present as graphitic carbon (and hence in the form of flakes) will not be affected by the graphitizing treatment and is undesirable in the final structure.

The temperatures used in malleabilizing usually range from about 1,500 to 1,650 °F. Time at temperature is quite long, frequently being as much as two to three days. Holding at this temperature causes the decomposition of the cementite into austenite plus rather rounded forms of graphite. If the iron is then cooled very slowly through the critical range (less than 10 °F/hr), the resultant structure will consist of ferrite plus temper carbon. Cooling less slowly through the transformation range, or using an iron containing an alloying element that retards graphitization, results in a final product containing various amounts of pearlite plus ferrite with, of course, temper carbon. Irons having a pearlitic structure plus temper carbon are known as pearlitic malleable irons (fig. 24, H). As might be expected, pearlitic malleable irons are susceptible to heat treatment in the same manner as steels.

7.3. Normalizing, Quenching, and Tempering

When heated to above the critical temperature (line SK, fig. 3), cast iron consists of austenite plus excess cementite or graphite and the austenite can be transformed in the same manner as in steel. Although irons that are almost completely graphitic (the common soft gray irons) can be hardened, this is not customary. Such irons must be held at temperatures well above the critical for long periods of time in order to assure the solution of carbon in the austenite.

For optimum results, the graphite flakes should be small and uniformly distributed and the matrix should be pearlitic with not too much free ferrite (combined carbon from about 0.50 to 0.80%). Castings should be heated uniformly to the quenching temperature, which usually ranges from 1,500 to 1,600 °F. The quenching medium commonly used is oil; water is used to a limited extent and some alloy castings may be quenched in air. The iron, as quenched, generally has a lower strength than as cast, but the strength is increased by tempering in the range of 350 to 1,200 °F. Softening proceeds uniformly as the tempering temperature is increased above about 350 to 1,200 °F.

A pearlitic matrix can be produced in nodular iron as cast or by normalizing. The normalizing treatment consists of heating to 1,600 to 1,650 °F, holding at temperature for solution of carbon, followed by cooling in air. The mechanical properties of the normalized irons vary with the composition, fineness, and amounts of pearlite; heavy sections are lower in hardness and higher in ductility than light sections. The Brinell hardness numbers range from about 200 to 275.

Martensitic or bainitic structures are produced in nodular iron by quenching from 1,600 to 1,700 °F in oil or water. The irons as quenched have high strengths but low ductility. However, the ductility of the as-quenched iron can be materially increased (strength and hardness decreased) by tempering at about 1,000 to 1,300 °F.

#### 7.4. Special Heat Treatments

Subject to the same limitations as steels, cast irons can be flame-hardened, induction-hardened, austempered, martempered, or nitrided.

#### 8. Practical Considerations

Satisfactory heat treatment of steel (and iron) requires furnaces that have uniform controlled temperatures, means for accurate temperature measurement, and for protecting the surface of the material from scaling or decarburizing. Proper quenching equipment is needed also.

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## 8.1. Furnaces and Salt Baths

There are many different types and sizes of furnaces used in heat treatment. As a general rule, furnaces are designed to operate in certain specific temperature ranges and attempted use in other ranges frequently results in work of inferior quality. In addition, using a furnace beyond its rated maximum temperature shortens its life and may necessitate costly and time-consuming repairs.

Fuel-fired furnaces (gas or oil) require air for proper combustion and an air compressor or blower is therefore a necessary adjunct. These furnaces are usually of the muffle type, that is, the combustion of the fuel takes place outside of and around the chamber in which the work is placed. If an open muffle is used, the furnace should be designed so as to prevent the direct impingement of flame on the work.

In furnaces heated by electricity the heating elements are generally in the form of wire or ribbon. Good design requires incorporation of additional heating elements at locations where maximum heat loss may be expected. Such furnaces commonly operate up to a maximum temperature of about 2,000 °F. Furnaces operating at temperatures up to about 2,500 °F usually employ resistor bars of sintered carbides.

Furnaces intended primarily for tempering may be heated by gas or electricity and are frequently equipped with a fan for circulating the hot air.

Salt baths are available for operating at either tempering or hardening temperatures. Depending on the composition of the salt bath, heating can be conducted at temperatures as low as 325 °F to as high as 2,450 °F. Lead baths can be used in the temperature range of 650 to 1,700 °F. The rate of heating in lead or salt baths is much faster than in furnaces.

#### a. Protective Atmospheres

It is often necessary or desirable to protect steel or cast iron from surface oxidation (scaling) and loss of carbon from the surface layers (decarburization, fig. 15, F). Commercial furnaces, therefore, are generally equipped with some means of atmosphere control. This usually is in the form of a burner for burning controlled amounts of gas and air and directing the products of combustion into the furnace muffle. Water vapor, a product of this combustion, is detrimental and many furnaces are equipped with a means for eliminating it. For furnaces not equipped with atmosphere control, a variety of external atmosphere generators are available. The gas so generated is piped into the furnace and one generator may supply several furnaces. If no method of atmosphere control is available, some degree of protection

may be secured by covering the work with cast iron borings or chips.

Since the work in salt or lead baths is surrounded by the liquid heating medium, the problem of preventing scaling or decarburization is simplified.

Vacuum furnaces also are used for annealing steels, especially when a bright non-oxidized surface is a prime consideration.

## b. Temperature Measurement and Control

Accurate temperature measurement is essential to good heat treating. The usual method is by means of thermocouples; the most common base-metal couples are copper-Constantan (up to about 700 °F), iron-Constantan (up to about 1,400 °F), and Chromel-Alumel (up to about 2,200 °F). The most common noblemetal couples (which can be used up to about 2,800 °F) are platinum coupled with either the alloy 87 percent platinum—13 percent rhodium or the alloy 90 percent platinum—10 percent rhodium. The temperatures quoted are for continuous operation.

The life of thermocouples is affected by the maximum temperature (which may frequently exceed those given above) and by the furnace atmosphere. Iron-Constantan is more suited for use in reducing and Chromel-Alumel in oxidizing atmospheres. Thermocouples are usually encased in metallic or ceramic tubes closed at the hot end to protect them from the furnace gases. A necessary adjunct is an instrument, such as a millivoltmeter or potentiometer, for measuring the electromotive force generated by the thermocouple. In the interest of accurate control, the hot junction of the thermocouple should be placed as close to the work as possible. The use of an automatic controller is valuable in controlling the temperature at the desired value.

If temperature-measuring equipment is not available, it becomes necessary to estimate temperatures by some other means. An inexpensive, yet fairly accurate method involves the use of commercial crayons, pellets, or paints that melt at various temperatures within the range 125 to 1,600 °F. The least accurate method of temperature estimation is by observation of the color of the hot hearth of the furnace or of the work. The heat colors observed are affected by many factors, such as the conditions of artificial or natural light, the character of the scale on the work, etc.

Steel begins to appear dull red at about 1,000 °F, and as the temperature increases the color changes gradually through various shades of red to orange, to yellow, and finally to white. A rough approximation of the correspondence between color and temperature is indicated in figure 25.

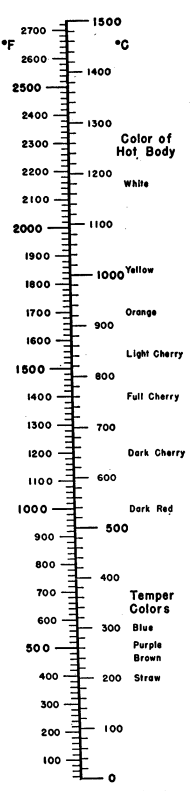


FIGURE 25. Temperature chart indicating conversion of Centrigrade to Fahrenheit or vice versa, color temperature scale for hardening-temperature range, and temperingtemperature range.

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It is also possible to secure some idea of the temperature of the piece of a carbon or lowalloy steel in the low temperature range used for tempering from the color of the thin oxide film that forms on the cleaned surface of the steel when heated in this range. The approximate temperature-color relationship for a time at temperature of about one-half hour is indicated on the lower portion of the scale in figure 25.

## 8.2. Quenching Media and Accessories

Quenching solutions act only through their ability to cool the steel. They have no beneficial chemical action on the quenched steel and in themselves impart no unusual properties. Most requirements for quenching media are met satisfactorily by water or aqueous solutions of inorganic salts such as table salt or caustic soda, or by some type of oil. The rate of cooling is relatively rapid during quenching in brine, somewhat less rapid in water, and slow in oil.

Brine usually is made of a 5- to 10-percent solution of salt (sodium chloride) in water. In addition to its greater cooling speed, brine has the ability to "throw" the scale from steel during quenching. The cooling ability of both water and brine, particularly water, is considerably affected by their temperature. Both should be kept cold—well below 60 °F. If the volume of steel being quenched tends to raise the temperature of the bath appreciably, the quenching bath should be cooled by adding ice or by some means of refrigeration.

There are many specially prepared quenching oils on the market; their cooling rates do not vary widely. A straight mineral oil with a Saybolt viscosity of about 100 at 100 °F is generally used. Unlike brine and water, the oils have the greatest cooling velocity at a slightly elevated temperature--about 100 to 140 °F—because of their decreased viscosity at these temperatures.

When steel is quenched, the liquid in immediate contact with the hot surface vaporizes; this vapor reduces the rate of heat abstraction markedly. Vigorous agitation of the steel or the use of a pressure spray quench is necessary to dislodge these vapor films and thus permit the desired rate of cooling.

Shallow hardening steels, such as plain carbon and certain varieties of alloy steels, have such a high critical cooling rate that they must be quenched in brine or water to effect hardening. In general, intricately shaped sections should not be made of shallow hardening steels because of the tendency of these steels to warp and crack during hardening. Such items should be made of deeper hardening steels capable of being hardened by quenching in oil or air.

A variety of different shapes and sizes of tongs for handling hot steels is necessary. It should be remembered that cooling of the area contacted by the tongs is retarded and that such areas may not harden, particularly if the steel being treated is very shallow hardening. Small parts may be wired together or quenched in baskets made of wire mesh.

Special quenching jigs and fixtures are frequently used to hold steels during quenching in a manner to restrain distortion.

When selective hardening is desired, portions of the steel may be protected by covering with alundum cement or some other insulating material. Selective hardening may be accomplished also by the use of water or oil jets designed to direct the quenching medium on the areas to be hardened. This also is accomplished by the induction and flame-hardening procedures previously described, particularly on large production jobs.

## 8.3. Relation of Design to Heat Treatment

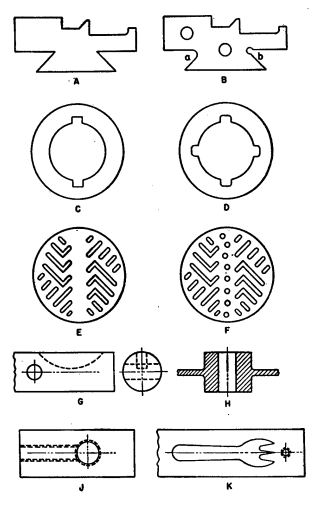
Internal strains arise from many causes, but the most serious are those developed during quenching by reason of differential cooling and from the increase in volume that accompanies the martensitic transformation. These stresses are frequently sufficient to distort or crack the hardened steel. Since temperature gradients are largely a function of the size and shape of the piece being quenched, the basic principle of good design is to plan shapes that will keep the temperature gradient throughout a piece at a minimum during quenching.

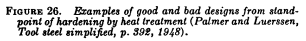
Because of the abruptness in the change of section, some shapes are impractical to harden, without cracking or distortion, by quenching in water, but a certain latitude in design is permissible when using an oil-hardening or airhardening steel. Other things being equal, temperature gradients are much lower in shapes quenched in oil than in water, and are still less in air. Thus a certain design may be perfectly safe for one type of steel, or one type of coolant, and unsafe for another.

Errors in design reach farther than merely affecting the internal strains during hardening. A sharp angle or notch serves to greatly concentrate the stresses applied during service, and the design of the part may be entirely responsible for concentrating the service stresses at a point already weakened by internal strains produced during hardening. Concentration of service stresses frequently parallels concentration of heat-treating strains and is frequently caused and cured by the same combination of circumstances.

A part is properly designed, from a standpoint of heat treatment, if the entire piece can be heated and cooled at approximately the same rate during the heat-treating operations.

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A, End view of an undercutting form tool incorrectly designed.

B, The same tool better designed from the viewpoint of heat treatment. Heavy sections have been lightened by drilling holes, thus insuring more uniform cooling. The fillet at (a) minimizes danger of cracking at the sharp recontrant angle. Where a fillet is not allowable, treatment as shown at (b) is helpful.

C. Cracking will tend to occur at the sharp roots of the keyways

C. Gracking will tend to occur at the sharp roots of the keyways.
D. Fillets at the roots of the keyways will reduce the tendency toward cracking. The incorporation of the two additional keyways, even though unnecessary in actual service, helps balance the section and avoid warping.
E. A blanking die with the center rib heavier than the surrounding areas. This may cause warping on quenching.
F. The same die with holes drilled in the center rib to equalize the amount of metal throughout the die, thus eliminating warpage difficulties.
G. A stem pinion with a keyway about one-half the diameter of the stem. The base of the keyway is extremely sharp, and the piece is further weakened by a hole drilled through the center of the stem nar the keyway. The base of the keyway should be filleted and hole re-located.
H. A dangerous design consisting of a thin collar adjoining a thick section. When hardening such pieces, the thin section often warps or cracks at the junction with the hub. Extremely generous fillets and drilling holes through the chalf.
J. When hardening, the concentration of strains at the junction of the two

J. When hardening, the concentration of strains at the junction of the two holes in the center are apt to cause failure. Such holes should be plugged before hardening.

K. A blanking die poorly designed. Crack will occur from point of fork prong to setscrew hole. The position of the setscrew hole should be changed to eliminate cracking.

Perfection in this regard is unattainable because, even in a sphere, the surface cools more rapidly than the interior. The designer should, however, attempt to shape parts so that they will heat and cool as uniformly as possible. The greater the temperature difference between any two points in a given part during quenching, and the closer these two points are together, the greater will be the internal strain and, therefore, the poorer the design.

When large and small sections are unavoidable in the same piece, the thick part frequently can be lightened by drilling holes through it. Where changes in section are encountered, angles should be filletted generously. Some examples of poor and good design are shown in figure 26.

#### 9. Nomenclature and Chemical **Compositions of Steels**

## 9.1. Structural Steels

In order to facilitate the discussion of steels, some familiarity with their nomenclature is desirable.

A numerical index, sponsored by the Society of Automotive Engineers (SAE) and the American Iron and Steel Institute (AISI), is used to identify the chemical compositions of the structural steels. In this system, a four-numeral series is used to designate the plain carbon and alloy steels; five numerals are used to designate certain types of alloy steels. The first two digits indicate the type of steel, the second digit also generally (but not always) gives the approximate amount of the major alloying element and the last two (or three) digits are intended to indicate the approximate middle of the carbon range. However, a deviation from the rule of indicating the carbon range is sometimes necessary.

The series designation and types are summarized as follows:

design	Types
10xx	 Nonsulphurized carbon steels
117X	 Resulphurized carbon steels (free ma-
	Rephosphorized and resulphurized carbon steels (free machining)
13xx	 Manganese 175%
<b>*23xx</b>	 Nickel 3.50%
TZ5XX	 Nickel 5.00%
31XX	 Nickel 1.25% chromium 0.65%
JOXX	 Nickel 3.50%, chromium 1 55%
40XX	 Molybdenum $0.20 \text{ or } 0.95\%$
41xx	 Chromium 0.50 or 0.95%, molybdenum
	0.12 or 0.20% Nickel 1.80%, chromium 0.50 or 0.80%, molybdenum 0.25%
44xx	 Molybdenum 0.40%
45XX	 Molybdenum 0 52%
46xx	 Nickel 1.80%, molybdenum 0.25%

- 47xx \_\_\_\_ Nickel 1.05%, chromium 0.45%, molybde-Nickel 1.05%, chromium 0.75%, m num 0.20 or 0.35% Nickel 3.50%, molybdenum 0.25% Chromium 0.25, 0.40 or 0.50% Carbon 1.00%, chromium 0.50% Chromium 0.80, 0.90, 0.95, or 1.00% 48xx \_\_
- 50xx \_\_\_\_
- 50xxx ....
- 51xx \_\_\_\_ 51xxx \_\_\_\_
- 51xxx ---- Carbon 1.00%, chromium 1.05% 52xxx ---- Carbon 1.00%, chromium 1.45% 61xx ---- Chromium 0.60, 0.80 or 0.95%, vanadium
- 0.12%, 0.10% min., or 0.15% min. 81xx \_\_\_\_ Nickel 0.30%, chromium 0.40%, molyb-denum 0.12%
- 86xx \_\_\_\_\_ Nickel 0.55%, chromium 0.50%, molyb-denum 0.20%
- 87xx \_\_\_\_ Nickel 0.55%, chromium 0.05%, molyb-denum 0.25%
- 88xx \_\_\_\_ Nickel 0.55%, chromium 0.50%, molyb-denum 0.35%
- 92xx \_\_\_\_ Manganese 0.85%, silicon 2.00%, chromium 0 or 0.35%
- 93xx \_\_\_\_ Nickel 3.25%, chromium 1.20%, molyb-denum 0.12%
- 94xx \_\_\_\_ Nickel 0.45%. chromium 0.40%, molybdenum 0.12%
- 98xx \_\_\_\_ Nickel 1.00%, chromium 0.80%, molyb-denum 0.25%

\* Not included in the current list of standard steels.

Listings of the AISI type numbers and chemical composition limits of the standard structural steels (complete as of January 1966) are given in table 1. The SAE type numbers (not shown in the table) are the same as the AISI numbers, except that the latter may be given a letter prefix to indicate the method of manufacture. These prefixes and their meanings are as follows: B denotes acid bessemer carbon steel, C denotes basic open hearth or basic electric furnace carbon steels, E denotes electric furnace alloy steels. A few of the AISI steels are not included in SAE listings.

Small quantities of certain elements are present in alloy steels that are not specified as These elements are considered as required. incidental and may be present to the maximum amounts as follows: copper, 0.35 percent; nickel, 0.25 percent; chromium, 0.20 percent; molybdenum, 0.06 percent.

The list of standard steels is altered from time to time to accommodate steels of proven merit and to provide for changes in the metallurgical and engineering requirements of industry.

Many of the alloy structural steels are manufactured to meet certain specified limits in hardenability as determined by the standard end-quench test. Such steels, table 2, are designated by the letter "H" following the AISI The chemical composition limits of number. these steels have been modified somewhat from the ranges or limits applicable to the same grades when specified by chemical composition only. The hardenability of an "H" steel is guaranteed by the manufacturer to fall within a hardenability band having maximum and minimum limits as shown by two limiting hardenability curves for that particular steel.

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Series

# TABLE 1. Composition limits of standard steels NONRESULPHURIZED CARBON STEELS •

Designation number	. Che	mical compositi	ion limits, perce	ent	Designation number	Chemical composition limits, percent					
AISI	C Mn		P(max) S(max		AISI	с	Mn	P(max)	S(max)		
*C 1005 *C 1006 C 1008 C 1010 *C 1011 C 1012	0.06 max .08 max .10 max 0.08/0.18 .08/0.18 .10/0.15	0.35 max 0.25/0.40 .25/0.50 .30/0.60 .60/0.90 .30/0.60	0.040 .040 .040 .040 .040 .040 .040	0.050 .050 .050 .050 .050 .050 .050	C 1042 C 1043 C 1044 C 1045 C 1046 C 1046 C 1048	0.40/0.47 .40/0.47 .43/0.50 .43/0.50 .43/0.50 .43/0.50 .44/0.52	0.60/0.90 .70/1.00 .80/0.60 .60/0.90 .70/1.00 1.10/1.40	0.040 .040 .040 .040 .040 .040 .040	0.050 .050 .055 .055 .055 .055		
*C 1013 C 1015 C 1016 C 1017 C 1017 C 1018 C 1019	.11/0.16 .13/0.18 .13/0.18 .15/0.20 .15/0.20 .15/0.20	.50/0.80 .30/0.60 .60/0.90 .30/0.60 .60/0.90 .70/1.00	.040 .040 .040 .040 .040 .040	.050 .050 .050 .050 .050 .050 .050	C 1049 C 1050 C 1051 C 1052 C 1053 C 1053 C 1055	.46/0.53 .48/0.55 .45/0.56 .47/0.56 .48/0.55 .50/0.60	.60/0.90 .60/0.90 .85/1.15 1.20/1.50 .70/1.00 .60/0.90	.040 .040 .040 .040 .040 .040 .040	.05( .05( .05( .05( .05( .05(		
C 1020 C 1021 C 1022 C 1028 C 1024 C 1025	.18/0.23 .18/0.23 .18/0.23 .20/0.25 .19/0.25 .22/0.28	.80/0.60 .60/0.90 .70/1.00 .80/0.60 1.85/1.65 0.80/0.60	.040 .040 .040 .040 .040 .040	.050 .050 .050 .050 .050 .050	*C 1059 C 1060 *C 1061 *C 1064 *C 1065 *C 1065 *C 1066	.55/0.65 .55/0.65 .55/0.65 .60/0.70 .60/0.70 .60/0.70	.50/0.80 .60/0.90 .75/1.05 .50/0.80 .60/0.90 .85/1.15	.040 .040 .040 .040 .040 .040	.05 .05 .05 .05 .05 .05		
C 1026 C 1027 C 1029 C 1030 *C 1084 C 1085	.22/0.28 .22/0.29 .25/0.31 .28/0.34 .32/0.38 .32/0.38	.60/0.90 1.20/1.50 0.60/0.90 .60/0.90 .50/0.80 .60/0.90	.040 .040 .040 .040 .040 .040	.050 .050 .050 .050 .050 .050 .050	*C 1069 C 1070 *C 1072 *C 1074 *C 1074 *C 1075 C 1078	.65/0.75 .65/0.75 .65/0.75 .70/0.80 .70/0.80 .72/0.85	.40/0.70 .60/0.90 1.00/1.30 .50/0.80 .40/0.70 .80/0.60	.040 .040 .040 .040 .040 .040 .040	.05 .05 .05 .05 .05 .05		
C 1086 C 1087 C 1088 C 1089 C 1040 C 1041	.30/0.37 .32/0.38 .35/0.42 .37/0.44 .37/0.44 .36/0.44	1.20/1.50 0.70/1.00 .60/0.90 .70/1.00 .60/0.90 1.35/1.65	.040 .040 .040 .040 .040 .040	.050 .050 .050 .050 .050 .050	C 1080 C 1084 *C 1086 C 1090 C 1095	.75/0.88 .80/0.93 .80/0.93 .85/0.98 .90/1.03	.60/0.90 .60/0.90 .30/0.50 .60/0.90 .80/0.50	.040 .040 .040 .040 .040 .040	.05 .05 .05 .05		

Designation number	Che	emical compositi	on limits, perce	ent	Designation number	Chemical composition limits, percent					
AISI	<b>c</b>	Mn	P(max)	S	AISI	с	Mn	P(max)	S		
C 1108 C 1109 C 1110 C 1116 C 1116 C 1117 C 1118 C 1119 C 1132	0.08/0.13 .08/0.13 .08/0.13 .14/0.20 .14/0.20 .14/0.20 .14/0.20 .14/0.20 .27/0.84	0.50/0.80 .60/0.90 .30/0.60 1.10/1.40 1.00/1.30 1.30/1.60 1.00/1.80 1.35/1.65	0.040 .040 .040 .040 .040 .040 .040 .04	0.08/0.13 .08/0.13 .08/0.13 .16/0.23 .08/0.13 .08/0.13 .24/0.33 .08/0.13	C 1137 C 1139 C 1140 C 1141 C 1144 C 1145 C 1145 C 1145 C 1146	0.32/0.39 35/0.43 37/0.44 37/0.45 40/0.48 42/0.49 42/0.49 42/0.49 48/0.55	1.35/1.65 1.35/1.65 .70/1.00 1.35/1.65 1.35/1.65 5.70/1.00 .70/1.00 .70/1.00	0.040 .040 .040 .040 .040 .040 .040 .04	0.08/0.13 .12/0.20 .08/0.13 .08/0.13 .24/0.33 .04/0.07 .08/0.13 .08/0.13		

ACID BESSEMER RESULPHURIZED CARBON STEELS •

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REPHOSPHORIZED AND RESULPHURIZED CARBON STEELS 4

Designation number	Che	mical compositi	ion limits, perce	ent	Designation number	Chemical composition limits, percent					
AISI	с	Mn	P	S	AISI	с	Mn	Р	S		
B 1111 B 1112 B 1113	0.13 max .18 max .18 max	0.60/0.90 .70/1.00 .70/1.00	0.07/0.12 .07/0.12 .07/0.12	0.08/0.15 .16/0.23 .24/0.88	C 1211 C 1212 C 1213 C 1213 C 1215 ‡C 12L14	0.13 max .18 max .18 max .09 max .15 max	0.60/0.90 .70/1.00 .70/1.00 .75/1.05 .80/1.20	0.07/0.12 .07/0.12 .07/0.12 .04/0.09 .04/0.09	0.08/0.15 .16/0.23 .24/0.33 .26/0.35 .25/0.35		

t Lead == 0.15/0.35 percent. See footnotes at end of table.

## TABLE 1. Composition limits of standard steels—Continued

OPEN HEARTH AND ELECTRIC FURNACE ALLOY STEELS .

V V
15           15           10           10           10           10           10           11           12           13           14           15
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TABLE 1. Composition limits of standard steels-Continued

OPEN HEARTH AND ELECTRIC FURNACE ALLOY STEELS - (continued)

Designation number	Chemical composition limits, percent											
AISI	с	Mn _	P(max)	S(max)	Si	Ni	Cr	Мо	v			
8720 *8785 8740 *8742	.18/0.23 .83/0.88 .38/0.43 .40/0.45	.70/0.90 .76/1.00 .75/1.00 .75/1.00	.085 .085 .085 .085 .085	.040 .040 .040 .040 .040	.20/0.85 .20/0.85 .20/0.85 .20/0.85	.40/0.70 .40/0.70 .40/0.70 .40/0.70	.40/0.60 .40/0.60 .40/0.60 .40/0.60	.20/0.30 .20/0.30 .20/0.30 .20/0.30 .20/0.30				
8822 9255 9260 *9262	.20/0.25 .50/0.60 .55/0.65 .55/0.65	.75/1.00 .70/0.95 .70/1.00 .75/1.00	.035 .035 .035 .035	.040 .040 .040 .040	.20/0.85 1.80/2.20 1.80/2.20 1.80/2.20	.40/0.70	.40/0.60	.30/0.40				
*E 9310 *9840 *9850	.08/0.13 .88/0.43 .48/0.53	.45/0.65 .70/0.90 .70/0.90	.025 .085 .085	.025 .040 .040	.20/0.85 .20/0.85 .20/0.85	8.00/8.50 .85/1.15 .85/1.15	.20/0.40 1.00/1.40 .70/0.90 .70/0.90	.08/0.15 .20/0.30 .20/0.30				

#### BORON STEELS .

#### These steels can be expected to have 0.0005 percent minimum boron content.

Designation number	Chemical composition limits, percent											
AISI	с	Mn	P(max)	S(max)	8i	Ni	Cr	Mo	v			
50B44 50B46 50B50 50B60	0.48/0.48 .48/0.50 .48/0.58 .55/0.65	0.75/1.00 .75/1.00 .75/1.00 .75/1.00 .75/1.00	0.035 .035 .085 .085 .085	0.040 .040 .040 .040 .040	0.20/0.85 .20/0.85 .20/0.85 .20/0.85 .20/0.85		0.40/0.60 .20/0.85 .40/0.60 .40/0.60					
51B60	.55/0.65	.75/1.00	.085	.040	.20/0.85		.70/0.90					
81B45	.48/0.48	.75/1.00	.085	.040	.20/0.85	0.20/0.40	.85/0.55	0.08/0.15				
94B17 94B30	.15/0.20 .28/0. <b>33</b>	.75/1.00 .75/1.00	.035 .085	.040 .040	.20/0.85 .20/0.85	.30/0.60 .30/0.60	.80/0.50 .80/0.50	.08/0.15 .08/0.15	` 			

Designation number									
AISI	C Mn		P(max) S(max)		Si Al		Cr	Мо	v
	0.88/0.48	0.50/0.70	0.085	0.040	0.20/0.40	0.95/1.30	1.40/1.80	0.30/0.40	

NITRIDING STEEL .

\* Standard steels for wire rods only.

Standard steel designations Up to C 1015 excl. Over C 1025 incl. per: When Silicon: When allicon is required, the following ranges and limits are commonly used:

The pho	osphorus	and	sulphur	limitations	for	each	steelmaking	process	are	as fo	llows:
1	Process							Perc	ent	maxii	num
								P			S
Basic e	lectric fi	Irnac	e							0	.025

Basic electric furnace0.025	0.025
Basic open hearth0.035	0.040
Acid electric furnace0.050	0.050
Acid open hearth0.050	0.050

Minimum silicon limit for acid open hearth or acid electric furnace alloy steel is 0.15 percent. Small quantities of certain elements are present in alloy steels which are not specified or required. These elements are considered as incidental and may, be present to the following maximum amounts: Copper, 0.35 percent; nickel 0.25 percent; chromium, 0.20 percent, and molybdenum, 0.65 percent. Where minimum and maximum sulphur content is shown it is indicative of resulphurized steels.

TABLE 2.	Standard	H-steels
OPEN HEARTH AND	ELECTRIC	FURNACE STEELS

number			Chemic	d composition limit	s, percent		
AISI	C	Mn	Si	Ni	Cr	Мо	v
1830 H	0.27/0.83	1 45 /0 05					
1885 H 1840 H	.82/0.38	1.45/2.05	0.20/0.85 .20/0.85				
1840 H	.87/0.44	1.45/2.05	.20/0.85		-		
			.20/0.30			•   • • • • • • • • • • • • • • • • • •	
8140 H	.37/0.44	0.60/1.00	.20/0.85	1.00/1.45	0.45/0.85		
8310 H	.07/0.18			· ·	0.40/0.00		
	.01/0.10	.80/0.70	.20/0.85	8.20/3.80	1.80/1.80		
4027 H	.24/0.30	60/1 00	90 /0 9E				
*4028 H 4087 H	.24/0.80	.60/1.00	.20/0.85 .20/0.85		• • • • • • • • • • • • • • • • • • • •	0.20/0.80	
4087 H   4047 H	.84/0.41	.60/1.00	.20/0.85			20/0.30	
	.44/0.51	.60/1.00 .60/1.00 .60/1.00 .60/1.00	.20/0.35 .20/0.35			20/0.80 .20/0.80 20/0.80	
4118 H 4180 H 4187 H 4187 H 4140 H 4142 H 4145 H	.17/0.23			}			
4180 H	27/0 88 1	.60/1.00 .80/0.70	.20/0.85		0.80/0.70	.08/0.15 .15/0.25 .15/0.25 .15/0.25 .15/0.25 .15/0.25	
4187 H	.84/0.41	.60/1.00	.20/0.85 .20/0.85		.75/1.20	.15/0.25	
4140 H	.87/0.44 .89/0.46	65/1 10	.20/0.85		.75/1.20	.15/0.25	
4142 H	.89/0.46	.65/1.10	.20/0.85		.75/1.20	.15/0.25	
4145 H 4147 H	.42/0.49	.65/1.10 .65/1.10 .65/1.10 .65/1.10 .65/1.10	.20/0.85		75/1 20	.15/0.25	*******
4150 H	.44/0.51 .47/0.54	.65/1.10	.20/0.85 .20/0.85		.75/1.20 .75/1.20	15/0.25	•••••
4161 H	.55/0.65	.65/1.10	.20/0.35		.75/1.20 .65/0.95	.15/0.25 .15/0.25	
1		.00/1.10	.20/0.85		.65/0.95	.25/0.85	
4820 H 4887 H	.17/0.23	.40/0.70	.20/0.85	1 55 /9 00		· ·	
4387 H	.84/0.41	.55/0.90 .55/0.90 .60/0.95	.20/0.85	1.55/2.00 1.55/2.00	.\$5/0.65	.20/0.80 .20/0.80 .20/0.80	
4840 H E4840 H	.87/0.44	.55/0.90	.20/0.85	1.65/2.00	.65/0.95 .65/0.95	.20/0.30	
E-104U II	.87/0.44	.60/0.95	.20/0.35	1.55/2.00	.65/0.95	.20/0.80	
4419 H	.17/0.23	AT /0				.20/0.00	
	•	.85/0.75	.20/0.35			.45/0.60	
4620 H	.17/0.23 .17/0.23 .23/0.29	.85/0.75	90 /0 95	· 1 FF /0 00		1	1
4621 H 4626 H	.17/0.28	.60/1.00	.20/0.85 .20/0.85	1.55/2.00 1.55/2.00 .65/1.05		.20/0.80	
4626 H	.23/0.29	.40/0.70	.20/0.35	65/1.05		.20/0.80	
4710 11	15/2 01		-			.15/0.25	
4718 H 4720 H	.15/0.21 .17/0.28	.60/0.95	.20/0.85	0.85/1.25	.80/0.60	.80/0.40	
	.11/0.20	.45/0.75	.20/0.85	0.85/1.25	.80/0.60	.15/0.25	
4815 H	.12/0.18	80/0 70	00 (0. 05				
4817 H	.14/0.20	.80/0.70 .30/0.70	.20/0.85 .20/0.85	3.20/3.80 3.20/3.80 3.20/3.80		.20/0.30	
4820 H	.17/0.28	.40/0.80	.20/0.35	3.20/3.80		.20/0.80 .20/0.80	
5120 H				0.20/0.00		.20/0.80	
5180 H	.17/0.23	.60/1.00	.20/0.85		.60/1.00		
5182 H	90 /0 95	.60/1.00 .50/0.90	.20/0.35		.60/1.00 .75/1.20 .65/1.10		
5185 H	82/0.88	.50/0.90	.20/0.35		.65/1.10		
5140 H	.32/0.38 .32/0.38 .37/0.44 .42/0.49	.60/1_00	.20/0.35 .20/0.35		.70/1.15		
5145 H	.42/0.49	.60/1.00 .60/1.05 .60/1.00	.20/0.35		.60/1.00		
5147 H 5150 H	.45/0.52	.60/1.05	.20/0.85		.60/1.00 .80/1.25		
5155 H	.47/0.54	.60/1.00	.20.0.35		.60/1.00		
5160 H	.50/0.60 .55/0.65	.60/1.00	.20/0.35		.60/1.00		
	.00/0.00	.65/1.10	.20/0.35		.60/1.00		
6118 H	.15/0.21	.40/0.80	.20/0.35				
6150 H	.47/0.54	.60/1.00	.20/0.35		.40/0.80		0.10/0
8617 H	14/0 00				.75/1.20		.15
8620 H	.14/0.20	.60/0.95 .60/0.95	.20/0.35 .20/0.35	0.85/0.75	.85/0.65	.15/0.25	1
8622 H	.17/0.23 .19/0.25	.60/0.95	.20/0.35	.85/0.75 .85/0.75	.35/0.65	.15/0.25	
8625 H I	.22/0.28	.60/0.95	.20/0.85	.85/0.75	.35/0.65 .35/0.65	.15/0.25	
8627 H	.24/0.80	.60/0.95	.20/0.35 .20/0.35 .20/0.35 .20/0.35	.85/0.75	.85/0.65	.15/0.25 .15/0.25	
8630 H	.27/0.83	.60/0.95	.20/0.35	.80/U.00 95/0.75	.85/0.65	.15/0.25	
8637 H	.84/0.41	.60/0.95 .70/1.05	.20/0.85	.35/0.75 .35/0.75 .35/0.65 .35/0.75 .35/0.75 .35/0.75	.85/0.65 .85/0.65	.15/0.25	
8640 H 8642 H	.87/0.44	.70/1.05	.20/0.35	.85/0.75	.85/0.65	.15/0.25	
8645 H	42/0 49	.70/1.05	20/0 85	.30/0.75	.85/0.65	15/0.25	
8655 H	.89/0.46 .42/0.49 .50/0.60	.70/1.05	.20/0.85 .20/0.85	.85/0.75	.85/0.65	.15/0.25	
			.20/0.85	.85/0.75	.85/0.65	.15/0.25 .15/0.25 .15/0.25 .15/0.25 .15/0.25	
8720 H	.17/0.28	.60/0.95	.20/0.85	.35/0.75	9F /A		
8740 H	.87/0.44	.70/1.05	.20/0.85	.85/0.75	.85/0.65 .85/0.65	.20/0.80	
8822 H	19/0 95				.00/0.65	.20/0.80	
0022 11	.19/0.25	.70/1.05	.20/0.35	.85/0.75	.85/0.65	.30/0.40	
9260 H	.55/0.65	.65/1.10	1.70/2.20		,		

#### **TABLE 2.** Standard H-steels—Continued

#### BORON H-STEELS

These steels can be expected to have 0.0005 percent minimum boron content

Designation number			Chemics	l composition limits,	percent		
AISI	с	Mn	Si	Ni	Cr	Mo	v
50B40 H 50B44 H 50B46 H 50B50 H 50B60 H	0.87/0.44 .42/0.49 .48/0.50 .47/0.54 .55/0.65	0.65/1.10 .65/1.10 .65/1.10 .65/1.10 .65/1.10	0.20/0.35 .20/0.35 .20/0.35 .20/0.35 .20/0.35 .20/0.35		0.80/0.70 .80/0.70 .18/0.48 .80/0.70 .80/0.70		
51B60 H	.55/0.65	.65/1.10	.20/0.85		.60/1.00		
81B45 H	.42/0.49	.70/1.05	.20/0.85	0.15/0.45	.30/0.60	0.08/0.15	
94B17 H 94B30 H	.14/0.20 .27/0.33	.70/1.05 .70/1.05	.20/0.85 .20/0.85	.25/0.65 .25/0.65	.25/0.55 .25/0.55	.08/0.15 .08/0.15	

\* Sulphur content 0.085/0.050 percent.

The phosphorus and sulphur limitations for each steelmaking process are as follows:

	P	S
Basic electric furnaceBasic open hearth		0.025
Acid electric furnace		0.050
Acid open hearth		0.050

Minimum silicon limit for acid open hearth or acid electric furnace alloy steel is 0.15 percent.

Small quantities of certain elements are present in alloy steels which are not specified or required. These elements are considered as incidental and may be present to the following maximum amounts: Copper, 0.35 percent; nickel, 0.25 percent; chromium, 0.20 percent; and molybdenum, 0.06 percent.

#### 9.2. Tool Steels

Technically, any steel used as a tool may be termed a tool steel. Practically, however, the term is restricted to steels of special composition that are usually melted in electric furnaces and manufactured expressly for certain types of services.

The current commonly used tool steels have been classified by the American Iron and Steel Institute into seven major groups and each commonly accepted group or subgroup has been assigned an alphabetical letter. Methods of quenching, applications, special characteristics, and steels for particular industries were considered in this type classification of tool steels which is as follows:

Group	Symbol and type
Water hardening W Shock resisting S	
Cold work{AN	il hardening Iedium alloy
Hot work H—(	ligh carbon—high chromium H 1 to H 19 incl. chromium base, H 20 to H 39 incl. tungsten base, H 40 to H 59
( 57 )	incl. molybdenum base)
High speed $ \begin{cases} T - T \\ M - M \end{cases}$	folybdenum base
Special purpose{FC}	ow alloy arbon tungsten

The AISI identification and type classification of tool steels is given in table 3. Each major group, identified by a letter symbol, may contain a number of individual types that are identified by a suffix number following the let-

#### TABLE 3. Identification and type classification of tool steels

Desig-			Identii	lying eler	nents, p	ercent			
AISI	c	Mn	8i	Cr	Ni	V	W	Mo	Co
			WATER	HARD	ENING	}			
W1	= 0.60/1.40						<b>.</b>		
W2 W4	• .60/1.40			0.25		0.25			
₩5	•.60/1.40 1.10			.50					
			SHOC	K RESI	TING				
51	0.50		•	1.50			2.50		
32	.50		1.00					0.50	
54	.55	0.80	2.00	]				.40	
85 86	.45	1.40	1.25	1.50				.40	
87	.50			3.25				1.40	
	· · · · ·	COLI	WORI	K; OIL	HARDE	ENING			
01	0.90	1.00		0.50			0.50		
02 06	.90	1.60	1.00					0.25	
07	1.45 1.20		1.00	.75			1.75		
	COLD	VORK;	MEDI	UM AL	LOY AI	R HAI	RDENIN	īG	•
A2	1.00			5.00				1.00	
A3	1.25			5.00				1.00	
A4	1.00	2.00		. 1.00		•	• •	1.00	
A5 A6	0.70	2.00		1.00				1.00	
Â7	2.25			5.25		4.75	6 1.00	1.00	
<b>A</b> 8	. 55			5.00			1.25	1.25	
A9	.50		·	5.00	1.50	1.00		- 1.40	
A10	1.35	1.80	1.25		. 1.80		-	1.50	1

See footnotes at end of table.

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ter symbol. The percentages of the elements shown in the table for each type are not to be considered as the mean of the composition

## TABLE 3. Identification and type classification of tool steels—Continued

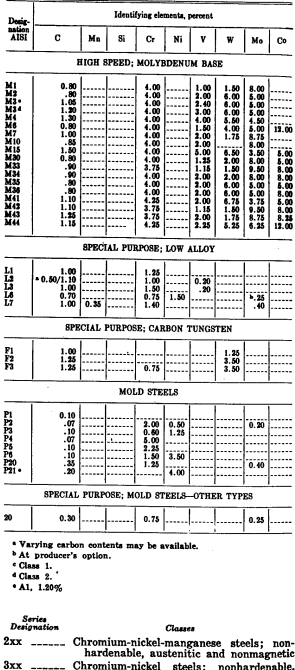
Desig-			Identi	fying ele	ments, j	percent			
nation AISI	С	Mn	8i	Cr	Ni	v	w	Mo	Co
H	COLD V	VORK;	HIGH (	CARBO	I N—HIG	H CHI	ROMIUI	l M	ł
D1 D2	1.00			12.00				1.00	
D3	1.50			12.00				1.00	
D4	2.25			12.00				1.00	
D5 D7	1.50			12.00				1.00	3.
D7	2.35			12.00		4.00		1.00	
		HOT	WORK;	CHRO	MIUM	BASE			·
H10	0.40			3.25		0.40		2.50	
H11 H12	.35	+		5.00		.40		1.50	
	.35			5.00 5.00		.40 1.00	1.50	1.50	
814	.40			5.00			5.00		
H16 H19	.55			7.00			7.00		
	. 40			4.25		2.00	4.25		4.
		HOT	WORK;	TUNG	sten :	BASE			
H20	0.35			2.00			9.00		
H21 H22	.35			3.50			9.00		
H23				2.00 12.00			11.00		
H24	.45			3.00			15.00		
H25 H26	.25			4.00			15.00		
a 20	.80			4.00	•••••	1.00	18.00		
	]	TOT W	ORK; N	OLYB	DENUM	I BASE	8		
141	0.65			4.00		1.00	1.50	8.00	
H42 H43	.60 .55			4.00 4.00		2.00	6 00	5.00	
				4.00		2.00		8.00	
		HIGH	SPEED	; TUNC	STEN	BASE			
[1]	0.70			4.00		1.00	18.00		
[2] [4]	.80	-		4.00		2.00	18.00		
5	.75	-		4.00		1.00 2.00	18.00 18.00		5.
r5 r6	. 80			4.50		1.50	20.00		8. 12.
[77	.75	-		4.00		2.00	14.00		
18 19	.75 1.20			4.00		2.00	14.00		5.
15	1.50	-		4.00		4.00	18.00 12.00		5.0
		-		1.00		0.00	14,00		. a.

ranges of the elements. Steels of the same type, manufactured by various producers, may differ in analysis from the values listed and may contain elements not listed in the type identification.

#### 9.3. Stainless and Heat Resisting Steels

The stainless and heat resisting steels possess relatively high resistance to oxidation and to attack by corrosive media at room and elevated temperatures. They are produced to cover wide ranges in mechanical and physical properties. They are melted exclusively by the electric furnace process.

A three-numeral system is used to identify stainless and heat resisting steels by type and according to four general classes. The first digit indicates the class and last two digits indicate type. Modification of types are indicated by suffix letters. The meaning of this AISI system is as follows:  
 TABLE 3. Identification and type classification of tool steels—Continued



S Desi	eries gnation	Classes
2xx		Chromium-nickel-manganese steels; non- hardenable, austenitic and nonmagnetic
3xx		Chromium-nickel steels; nonhardenable, austenitic and nonmagnetic
4xx		Chromium steels; hardenable, martensitic and magnetic
4xx		Chromium steels; nonhardenable, ferritic and magnetic
5xx		Chromium steels; low chromium heat re- sisting.

The chemical composition ranges and limits of the standard stainless and heat resisting steels are given in table 4.

							Chemical con	Chemical composition limits, percent	, percent					ľ	
AISI	0	(max) M	P (max)	S (max)	Si (max)	ō	ÿ	Mo	రి	æ	ä	Cb-Ta	£	V	N
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	Ne na	<b>b5.60/ 7.50</b>	0000 0000	0.030	88	16.00/18.00 17.00/19.00	3.50/ 5.50								0.25 max .25 max
301 302 302 303 303 803 803 803 803 803 803 803 803	X C C C C C C C C C C C C C C C C C C C	44444 88888	33388	.030 .030 .030 .030 .030 .030 .030	• 2.00/3.00 1.00 1.00 1.00	16.00/18.00 17.00/19.00 17.00/19.00 17.00/19.00 17.00/19.00	6.00/ 8.00 8.00/10.00 8.00/10.00 8.00/10.00 8/00.10.00	• 0.60 mex		0.16 min					
2004L 2004L	.08 max 03 max 12 max 0.03 max 0.03 max	44444 88888	.045 045 045 045 045	888 888 888 888 888 888 888 888 888 88	88888	18.00/20.00 18.00/20.00 17.00/19.00 19.00/21.00 22.00/24.00	8.00/12.00 8.00/12.00 10.00/13.00 12.00/15.00								
3005 310 314 316 314	.08 max 25 max 08 max 25 max 26 max	4444 88888 88888	349. 349. 349. 349. 349.	88888	1.00 1.50 1.50 1.50 1.00 1.00	22.00/24.00 24.00/26.00 24.00/26.00 23.00/26.00 16.00/18.00		<u></u>							
316L 317 321 347	20.00 20.000	44444 88888 88888		888888	88888	16.00/18.00 18.00/20.00 17.00/19.00 17.00/19.00	10.00/14.00 11.00/15.00 9.00/13.00 9.00/13.00 9.00/13.00	2.00/3.00 3.00/4.00	0.20 max		δ x C min	10 × C His 10 × C His 10 × C His	0. 10 max		
408 14 14 16 14 16	16 max 80. 16 max 11. 15 max 12.		940 940 940 940 940 940 940 940 940 940	.030 .030 .030 .030 .030	88888 88888	11.50/13.00 11.50/14.50 11.50/13.50 11.50/13.50 11.50/14.00	1.26/ 2.50	* 0.60 max						0.10/0.30	
416 Se 420 430 430F Se 430F Se	. 15 max Over 0. 15 0. 12 max . 12 max	1.28	88888	.060 .030 .030 .030 .030	88888	12.00/14.00 12.00/14.00 14.00/18.00 14.00/18.00 14.00/18.00		*.60 mex		0. 10 mm 0. 10 mm . 15 min					
431 440A 440B 440C 442	.20 max .60/0.75 .75/0.95 .95/1.20 .20 max	88888	555555	8888888	888888	15.00/17.00 16.00/18.00 16.00/18.00 18.00/18.00 18.00/18.00 23.00/27.00	1.25/2.50	75 max 75 max 75 max							26 max
501 502	Over 0.10 0.10 max	88.1	9 <del>0</del> .	030		4.00/ 6.00 4.00/ 6.00		. 40/0.65							

At producer's option: reported only when intentionally added. b Range.

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## 10. Recommended Heat Treatments

#### 10.1. Structural Steels

A listing of recommended heat treatments for many of the steels listed in tables 1 and 2 are given in tables 5 and 6. Treatments for the standard steels that are omitted can be estimated by interpolation (also see fig. 11). Where different quenching media are suggested, the smaller sections may harden satisfactorily in the slower quenching medium. The upper end of the heat treating ranges should always be used for large size sections. The inclusion of normalizing and annealing temperatures does not imply that these treatments are necessarily a prelude to hardening.

For hardening, the general rule is to place the cold steel in a furnace containing a slightly oxidizing atmosphere or neutral salt bath at the hardening temperature and let it heat naturally, hold ½ hr/in of thickness, and then quench; it is sometimes advisable to warm intricate sections or complex shapes before putting them into the high temperature unit and to remove from the quenching medium while still warm. Usually, the quenched piece should be allowed to cool until it feels comfortable to the bare hand and then immediately tempered. To minimize cracking, it is sometimes desirable to temper the quenched piece before it reaches room temperature; a double temper (i.e., repeating the treatment) is recommended for this condition.

For carburizing, most of the steels may be given either (1) a single, (2) double, or (3) triple treatment as is indicated in table 5. The time at the carburizing temperature will vary depending upon the temperature used and the desired thickness of the case. The relation between time, temperature and case depth is illustrated in figure 19.

AISI or SAE number	Carburizing temperature	Cooling method	Reheat	Cooling medium	2d reheat	Cooling medium	Tempering temperature
0.14.1004.11004.1445	• #	(Brine or water	• 7		°F		• F
003 to 1024; 1108 to 1119; 1211 to 1215	1 050 /0 500	Water or oil	1,400/1,450	Brine or water			1
40 1210	1,650/1,700	Cool slowly	1.400/1.450	dodo			250/3
		{do	1,650/1,700	Water or oil	1,400/1,450	Brine or water	J
		(011					h
810	1,650/1,700	do	1.875/1.425	Oil			250/3
		Cool slowly	1.875/1.425	do			1 200/0
		(do	1,52\$/1,575	do	1,375/1,425	Oil	l)
019 += 4094. 4119		Oildo	1,425/1,475				h
012 to 4024; 4118	1,650/1,700	Cool slowly	1,425/1,475	Oil	1		250/3
		{do	1,600/1,650	do	1,425/1,475	Oil	
		(Oii					ľ.
419,4422 and 4427; 8822	1,650/1,700	do	1,450/1,500	Oil			
	1,000/1,100	Cool mowly	1.450/1.500	do			} 250/8
		{do	1,625/1,675	do	1,450/1,500	Oil	IJ
		(011	;				h
520	1,650/1,700	do	1,475/1,525 1,475/1,525	Oil			11 050 /
•		Cool slowly	1,475/1,525	do			} 250/800
		(do	1,650/1,700	do	1,475/1,525	Oil	IJ
820; 4615 to 4626; 4718 and		Oildo					h
4720	1,650/1,700	Cool slowly	1,425/1,475	Oildo			250/
		dodo	1,525/1,575	do	1 875/1 425	Oil	
		(Oil			1,010/1,420	04	
815 to 4820	1,650/1,700	do	1.850/1.400	Oil			
	-,000/1,700	Cool slowly	1.850/1.400	0			250/
		(do	1,550/1,600	do	1,850/1,400	Oil	-U
		(Oil					.h
015; 5115 and 5120	1,650/1,700	do	1,425/1,475	l Oil			250/
		Cool slowly	1,425/1,475 1,600/1,650	do		Oil	·]( <sup>200/</sup>
		•	1,000/1,000	uo	1,420/1,475	UI	·μ
118 and 6120	1 050 /1 500	(Oildo	1.450/1.500	Oil or water			-h
110 anu 0140	1,650/1,700	Cool slowly	1 450/1 500	dodo			- 800/400
		(do	1,600/1,650	Oil	1,450/1,500	Oil or water	: ] 
		(Oil					Ľ.
115; 8615 to 8622; 8720; 8822;	1,650/1,700	}do	1.450/1.500	Oil			·[]
9310 and 94B17.	1,000/1,100	Cool alowly	1,450/1,500 1,450/1,500	do			250/3
		do	1,550/1,600	do	1.450/1.500	Oil	11

TABLE 5. Recommended heat treatments for carburizing grade ste	TABLE 5.	E 5. Recommended h	eat treatments	for carburizing	i arade steel
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TABLE 6.	Recommended	heat	treatments	for	heat	treati	ng	grade	steel	18
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AISI or SAE number	Normalizing temperature	Annealing temperature	Hardening temperature	Quenching medium	AISI or SAE number	Normalizing temperature	Annealing temperature	Hardening temperature	Quenching medium
1030 1040 1050 1060 1070 1080 1190 1152 1140 1151 1151 1151 11340 3140 4028 4042	°F 1,625/1,725 1,600/1,700 1,550/1,650 1,500/1,600 1,500/1,600 1,500/1,600 1,475/1,575 1,475/1,575 1,625/1,725 1,600/1,700 1,550/1,650 1,650/1,650 1,650/1,650 1,650/1,650	°F 1,525/1,575 1,475/1,525 1,450/1,525 1,425/1,475 1,425/1,475 1,375/1,425 1,375/1,425 1,525/1,575 1,450/1,525 1,450/1,525 1,475/1,525 1,475/1,525	°F 1,550/1,600 1,500/1,550 1,475/1,525 1,450/1,550 1,450/1,500 1,460/1,450 1,400/1,450 1,550/1,600 1,575 1,525/1,575 1,550/1,650 1,475/1,525 1,550/1,600	Water or brine. Do. Do. Do. Do. Do. Do. Do. Oil or water. Do. Oil. Oil or water. Oil.	5046 5130 5145 5160 5100 51100 52100 6150 8630 8630 8645 8735 8735 8735 8735 8745 9260 9840 9840	*F 1.600/1.700 1	*F 1.475/1.525 1.500/1.550 1.475/1.525 1.450/1.550 1.460/1.450 1.400/1.450 1.500/1.500 1.500/1.550 1.500/1.500/1.500 1.500/1.500/1.500 1.500/1.500	°F 1,500/1,550 1,550/1,600 1,525/1,575 1,600/1,550 1,450/1,550 1,450/1,550 1,550/1,600 1,525/1,575 1,525/1,575 1,525/1,575 1,525/1,575 1,550/1,600 1,500/1,550	Oil. Oil or water. Oil. Do. Do. Do. Do. Do. Do. Do. Do. Do. Do
4063 4130 4140 4150 4340	1,550/1,650 1,600/1,700 1,600/1,700 1,600/1,700 1,600/1,700	1,450/1,500 1,525/1,575 1,500/1,550 1,475/1,525 1,500/1,550	$\begin{array}{c} 1,475/1,526\\ 1,550/1,600\\ 1,525/1,575\\ 1,500/1,550\\ 1,500/1,550\\ 1,500/1,550\end{array}$	Do. Oil or water. Oil. Do. Do.	3000	1,600/1,700	1,475/1,525		

(No tempering treatments are given, as these temperatures depend upon the desired hardness)

#### 10.2. Tool Steels

Recommended heat treatments for the different types of tool steels are summarized in table 7.

The time the steel is at the normalizing temperature, after being heated uniformly through, varies from about 15 min for a small section to about 1 hr for large sizes; cooling from the normalizing temperature is in still air. For annealing, the upper range in temperature is recommended for large and the lower limit for small sections. The time the steel is at the annealing temperature, after being heated uniformly through, ranges from about 1 hr for thin sections of carbon or low alloy to about 4 hr for heavy sections of high alloy steel. Cooling from the annealing temperature through the transformation range should not exceed about 75 °F/hr for the carbon and low alloy and 50 °F/hr for the high alloy steels.

For hardening, the protective atmosphere, rate of heating, hardening temperature, time at temperature, and rate of cooling vary with the composition of the steels. A general rule is to heat the plain carbon and low alloy steels rapidly in a furnace having a slightly oxidizing atmosphere or in a neutral salt bath at the hardening temperature, hold 1/2 hr/in of thickness, and ther quench. The high alloy steels, such as the T and M types, are usually preheated to minimize distortion and cracking and then heated rapidly in a furnace containing a highly carburizing atmosphere or in a molten bath at the hardening temperature, held at temperature for time as indicated in the table, and then quenched. Tools should be tempered immediately after quenching; a double temper is recommended for the high alloy steels.

The carburizing steels (type P) may be (1) quenched directly from the carburizing temperature; (2) quenched directly from the carburizing temperature followed by reheating to and quenching from the recommended hardening temperature; (3) slowly cooled from the carburizing temperature followed by reheating to and quenching from the recommended hardening temperature or (4) slowly cooled from the carburizing temperature, reheated to the carburizing temperature and quenched, followed by reheating to and quenching from the recommended hardening temperature. For most purposes, method (3) is generally suitable. Tempering should follow in all cases.

#### 10.3. Stainless and Heat Resisting Steels

The response of stainless and heat resisting steels to heat treatment depends upon their composition. The general methods employed are similar to those used with other steels and no special equipment is needed. Heat treatment may be performed in any conventional electric, gas fired, oil fired, salt bath, or induction furnace in which the required temperatures can be obtained. Precautions should be taken to avoid exposure of the steel to the direct flame of the burners. Conditions that will result in carburization of the surface of the steel should be avoided. In general, oxidizing furnace atmospheres are preferred. Bright, scale-free heat treatment is possible by the use of specially prepared and controlled atmospheres.

TABLE 7. Recommended heat treatments for tool steels

AISI	Normalizing	Anner	ling		H	lardening	Tempering		
type	temperature	Temperature	Brinell hardness number	Preheating temperature	Quenching temperature	Quenching medium	Temperature	Rockweil hardness number	
W1 W2 W4 W5	•F • 1,450/1,700 • 1,450/1,700 • 1,450/1,700 • 1,450/1,700	°F 1,375/1,450 1,400/1,450 1,375/1,450 1,400/1,450	(BHN) 159/202 159/202 159/202 163/202	• <b>F</b> (0) (0)	°F 1,400/1,500 1,400/1,500 1,400/1,500 1,400/1,500	Brine or water do do	°F 850/650 850/650 850/650 850/650	(Rc) 64/( 64/( 64/( 64/(	
81 82 84 85 86	000000000000000000000000000000000000000	1,450/1,500 1,400/1,450 1,400/1,450 1,400/1,450	183/229 192/217 192/229 192/229 192/229	(b) 1,100/1,200 1,100/1,200 1,800/1,400		Oil Brine or water Oil do Air or oil	400/1,200 300/800 350/800	58/4 60/8 60/8	
87 01	(*) (*) (*) 1,500/1,600	1,475/1,525 1,500/1,550 1,400/1,450	187/228	1,400 1,200/1,800	1,675/1,750			60/1 56/1 57/4	
02 06 07	1,500/1,550 1,600/1,650 1,600/1,650	1,875/1,425 1,400/1,450 1,450/1,500	183/212 183/212 183/217 192/217	1,100/1,200 1,100/1,200 1,100/1,200 1,100/1,200	1,450/1,500 41,400/1,475 41,450/1,500 1,450/1,525 1,550/1,625	Oildo do Water Oil	800/500 300/500 800/600 825/550	62/1 62/1 63/1 64/1	
A2 A8 A4 A5 A6 A7 A8 A9 A10	0 0 0 0 1,450	$\begin{array}{c} 1,550/1,600\\ 1,550/1,600\\ 1,350/1,400\\ 1,350/1,400\\ 1,350/1,375\\ 1,600/1,550\\ 1,550/1,600\\ 1,550/1,600\\ 1,410/1,460\\ \end{array}$	202/229 207/229 200/241 228/255 217/248 235/262 192/223 212/248 235/269	1,850/1,450 1,450 1,150/1,250 1,100/1,200 1,100/1,200 1,400/1,500 1,450 1,450 1,200	1,700/1,800	Airdo do do do do do do	\$50/1,000 \$50/1,000 \$00/800 \$00/800 \$00/1,000 \$50/1,100 \$50/1,100 \$50/1,100	62/5 65/5 60/5 60/5 67/5 56/5 56/8 62/5	
01 02 08 04 05 07		1,600/1,650 1,600/1,650 1,600/1,650 1,600/1,650 1,600/1,650 1,600/1,650	207/248 217/255 217/255 217/255 223/255 235/262	1,400/1,500 1,400/1,500 1,400/1,500 1,400/1,500 1,400/1,500 1,400/1,500	1,775/1,850 1,800/1,875 1,700/1,800 1,775/1,850 1,800/1,875 1,850/1,950	Air Oil Air dodo	400/1,000 400/1,000 400/1,000 400/1,000 400/1,000	61/5 61/5 61/5 61/5 61/5 61/5 65/5	
110 111 112 113 114 114 119 220 221 23 24 25 26 26 26 41 42 43	000000000000000000000000000000000000000	$\begin{array}{c} 1,550/1,650\\ 1,550/1,650\\ 1,550/1,650\\ 1,550/1,650\\ 1,500/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,500/1,500\\ 1,500/1,500$	192/229 192/229 192/229 192/229 207/285 207/285 207/285 207/285 213/241 207/285 213/241 207/285 217/241 207/285 207/235 207/235	$\begin{array}{c} 1,500\\ 1,400/1,500\\ 1,400/1,500\\ 1,400/1,500\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,650\\ 1,350/1,550\\ 1,350/1,550\\ 1,350/1,550\\ \end{array}$	$\begin{array}{c} 1,850/1,900\\ 1,825/1,875\\ 1,825/1,875\\ 1,825/1,900\\ 1,850/1,950\\ 2,050/2,150\\ 2,000/2,200\\ 2,000/2,200\\ 2,000/2,200\\ 2,000/2,200\\ 2,000/2,200\\ 2,000/2,200\\ 2,000/2,200\\ 2,000/2,250\\ 2,000/2,250\\ 2,150/2,300\\ 2,150/2,300\\ 4,2,000/2,175\\ 4,2,000/2,175\\ 4,2,000/2,175\\ 4,2,000/2,175\\ 5,000/2,100\\ 5,000/2,100\\ 5,000/2,100\\ 5,000/2,100\\ 5,000/2,100\\ 5,000/2,100\\ 5,000/2,100\\ 5,000/2,100\\ 5,000/2,100\\ 5,000/2,100\\ 5,000/2,100\\ 5,000/2,100\\ 5,000/2,100\\ 5,000/2,100\\ 5,000/2,100\\ 5,0$	Air do dodo Air or oil do do do do do do do do do Air, oil, or salt do Air, oil, or salt do	1.000/1.200 1.000/1.200 1.000/1.200 1.000/1.200 1.000/1.200 1.000/1.200 1.000/1.250 1.000/1.250 1.000/1.250 1.100/1.250 1.000/1.250 1.050/1.250 1.050/1.250 1.050/1.250 1.050/1.250 1.050/1.200 1.050/1.200	56/3 56/3 55/3 55/3 55/3 55/3 55/3 47/4 56/4 56/4 56/5 55/4 56/5 56/5 56/5 55/4	
1 2 4 5 5 6 7 7 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	00000000	$\begin{array}{c} 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ \end{array}$	217/255 223/255 229/269 235/275 248/293 217/255 229/255 235/277 241/277	$\begin{array}{c} 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\end{array}$	• 2,300/2,375 • 2,300/2,375 • 2,300/2,375 • 2,325/2,400 • 2,325/2,400 • 2,325/2,400 • 2,300/2,350 • 2,300/2,375 • 2,275/2,325 • 2,200/2,300	Oil, air, or sait do do do do do do do do do	f 1,000/1,100 f 1,000/1,100 f 1,000/1,100 f 1,000/1,100 f 1,000/1,100 f 1,000/1,100 f 1,000/1,100 f 1,000/1,100 f 1,000/1,200	65/6 66/6 65/6 65/6 65/6 65/6 65/6 65/6	
1 3 Class 1 3 Class 1 4 4 6 7 10 15 80 83 84 85 86 41 42 43 44	000000000000000000000000000000000000000	$\begin{array}{c} 1,500/1,600\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,600/1,650\\ 1,500/1,600\\ 1,500/1,600\\ 1,500/1,600\\ 1,600/1,650\\ 1,600/1,600\\ 1,600/1,600\\ 1,600/1,600\\ 1,600/1,600\\ 1,600/1,600\\ 1,600/1,600\\ 1,600/1,600\\ 1,600/1,600\\ 1,600/1,600$	207/235 212/241 223/255 223/255 223/255 248/277 217/255 207/235 241/277 235/269 235/269 235/269 235/269 235/269 235/269 235/269 235/269 235/269 235/269 235/269 235/269 235/269 235/269	$\begin{array}{c} 1,350/1,550\\ 1,350/1,550\\ 1,350/1,500\\ 1,350/1,500\\ 1,350/1,500\\ 1,350/1,500\\ 1,350/1,550\\ 1,500/1,600\\ 1,350/1,550$	$\begin{array}{c} \cdot 2,150/2,225\\ \cdot 2,175/2,250\\ - 2,200/2,250\\ - 2,200/2,250\\ \cdot 2,200/2,250\\ \cdot 2,200/2,250\\ \cdot 2,150/2,200\\ \cdot 2,150/2,200\\ - 2,150/2,220\\ - 2,150/2,250\\ - 2,200/2,200\\ - 2,200/2,200\\ - 2,200/2,200\\ - 2,200/2,200\\ - 2,200/2,200\\ - 2,200/2,200\\ - 2,200/2,200\\ - 2,200/2,200\\ - 2,200/2,200\\ - 2,200/2,200\\ - 2,200/2,200\\ - 2,200/2,200\\ - 2,200/2,200\\ - 2,200/2,200\\ - 2,200/2,200\\$	Oil, sir, or salt do	1,000/1,100 1,000/1,100 1,000/1,100 1,000/1,100 1,000/1,100 1,000/1,100 1,000/1,100 1,000/1,100 1,000/1,100 1,000/1,100 1,000/1,100 1,000/1,100 1,000/1,100 1,000/1,100 1,000/1,100 1,000/1,100 1,000/1,100 1,000/1,100	85/65 65/66 65/61 66/61 66/61 66/61 66/61 65/65 65/65 65/65 65/65 65/65 65/65 66/60 66/60 66/60 66/60	
	1,600/1,650	1,425/1,475 1,400/1,450	179/207 163/196	(b) (b)	1,450/1,550 1,450/1,550 1,550/1,700	Oil or water Water Oil	800/600 800/1,000	64/54 63/4	
	1,600/1,650	1,450/1,500 1,400/1,450	174/201 183/212	(b) (b)	1,450/1,550 1,450/1,550 1,550/1,700 1,425/1,500 1,500/1,600 1,450/1,550 1,500/1,600	Oil Water Oil do	800/600	64/5	
		1,450/1,500 1,400/1,475 1,450/1,500 1,450/1,500	183/212 183/207 207/235	1,200		Water or brine	300/1,000 300/600 350/500	62/4 64/5 64/6	
	1,650	1,450/1,500	212/248	1,200 1,200	1,450/1,600	Water, brine, or oil	850/500 350/500 350/500	64/0 65/0 65/0	

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## TABLE 7. Recommended heat treatments for tool steels-Continued CARBURIZING GRADES

AISI	Normalizing	Annea	ling	Carburizing Harde		ardening	Tempering	
type	temperature	Temperature	Hardness	temperature	Temperature	Quenching medium	Temperature	Rc of case
P1 P2 P3 P4 P5 P6 P20 P21	• F (e) (e) (c) (e) (e) (e) (e) 1,650	°F 1,350/1,650 1,350/1,500 1,350/1,500 1,650/1,600 1,550/1,600 1,400/1,450 ( <sup>h</sup> )	(BHN) 81/100 103/123 109/137 116/128 105/110 190/210 150/180	°F 1,650/1,700 1,650/1,700 1,775/1,825 1,650/1,700 1,650/1,700 1,650/1,700	°F 1,450/1,475 1,525/1,550 1,475/1,525 1,775/1,825 1,550/1,600 1,450/1,500 1,500/1,600 i1,300/1,500	Water or brine           Oildo           Air           Oil or water           Oil           Air           Oil or water	°F 350/500 350/500 350/500 350/500 350/450 350/450 350/500 j 950/1,025	64/58 64/58 64/58 64/58 64/58 61/58 61/58 64/58 64/58
P21         1,050         (*)           • Normalizing:         0.60 to 0.75% C, 1500 °F;         .75 to .90% C, 1450 °F;           .90 to 1.10% C, 1600 °F;         .10 to 1.40% C, 1600 °F;           1.10 to 1.40% C, 1600 °to 1700 °F.           Annealing:           0.60 to 0.90% C, 1360 ° to 1400 °F           .90 to 1.40% C, 1400 ° to 1450 °F.					<sup>c</sup> Do not no <sup>d</sup> Do not so <sup>e</sup> Time at l <sup>f</sup> Double te <sup>g</sup> Normalisi <sup>b</sup> Do not an <sup>l</sup> Solution f	oak. hardening temperature, mpering recommended. ng not required.		

TABLE 8.	Recommended heat treatments for stainless and heat resisting steels
GROUP	I. HARDENABLE CHROMIUM STEELS (MARTENSITIC AND MAGNETIC)

AISI	Anneal	ing *	Harden	Hardening <sup>b</sup> Stress relieving		ieving	Tempering		
type	Temperature	Hardness	Temperature	Hardness	Temperature	Hardness	Temperature	Hardness	
403 410 414 416 416 Se 420 431 440 A	°F 1,550/1,650 1,500/1,650 • 1,200/1,300 1,550/1,650 1,550/1,650 • 1,550/1,650 • 1,150/1,225 1,625/1,675	75/83 R <sub>B</sub> 75/85 R <sub>B</sub> 98 R <sub>8</sub> /24 R <sub>e</sub> 78/83 R <sub>B</sub> 80/90 R <sub>B</sub> 81/89 R <sub>B</sub> 97 R <sub>B</sub> /24 R <sub>e</sub> 91/95 R <sub>B</sub>	°F 1,700/1,850 1,800/1,900 1,700/1,850 1,700/1,850 1,800/1,900 1,800/1,900 1,800/1,950	39/43 R. 39/43 R. 42/47 R. 40/44 R. 39/43 R. 53/56 R. 42/46 R. 55/58 R.	°F 450/700 450/700 450/700 450/700 300/700 300/700 300/700	40/37 R. 40/37 R. 44/40 R. 40/37 R. 53/48 R. 42/36 R. 56/51 R.	°F 1,000/1,100 1,000/1,400 1,100/1,800 1,000/1,400 1,000/1,400	34 R <sub>e</sub> /86 R 34 R <sub>e</sub> /86 R 29 R <sub>e</sub> /99 R 34 R <sub>e</sub> /86 R 34 R <sub>e</sub> /86 R 35 R <sub>e</sub> /99 R	
440 B 440 C 501 502	1,625/1,675 1,625/1,675 1,525/1,600 1,525/1,600	94/98 R <sub>B</sub> 95/99 R <sub>B</sub> 80/90 R <sub>B</sub> 72/80 R <sub>B</sub>	1,850/1,950 1,850/1,950 1,600/1,700 ( <sup>4</sup> )	57/59 R <sub>o</sub> 60/62 R <sub>o</sub> 35/45 R <sub>e</sub>	800/700 300/700	58/53 R <sub>c</sub> 60/55 R <sub>c</sub>	400/1,200	45/25 R	

# GROUP II. NONHARDENABLE CHROMIUM STEELS (FERRITIC AND MAGNETIC)

	Annealing •					
AISI type	Temperature	Hardness, R <sub>B</sub>				
	۰F					
105	1,850/1,500	70/80				
180	1,400/1,500	80/90				
180 F	1.850/1.500	80/90				
430 F Se	1.850/1.500	80/90				
142	1,800	80/90				
146	1.450/1.600	80/90				

# GROUP III. NONHARDENABLE CHROMIUM-NICKEL AND CHROMIUM-NICKEL-MANGANESE STEELS (AUSTENITIC AND NONMAGNETIC)

AISI	Annealing	AISI	Annealing
type	temperature '	type	temperature '
201 202 301 302 B 303 Se 303 Se 304 304 304 305 305 308	°F 1,850/2,050 1,850/2,050 1,850/2,050 1,850/2,050 1,850/2,050 1,850/2,050 1,850/2,050 1,850/2,050 1,850/2,050 1,850/2,050 1,850/2,050	309 309 S 810 810 S 314 316 L 817 821 847 848	°F 1,960/2,050 1,900/2,050 1,900/2,100 1,900/2,100 1,850/2,050 1,850/2,050 1,850/2,050 1,850/2,050 1,850/2,050 1,850/2,050

Cool slowly to 1,100 °F.
Although these steels harden appreciably on air cooling, quenching in oil is preferable.
Full annealing impractical; may be air cooled from indicated temperature.
Generally used in annealed condition only.
Air cool.
Cooling may be in air or by quenching in water. The resultant hardness of all of these steels will be in the range of approximately Rockwell B 80 to 90.

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The stainless steels may be divided into three general groups as follows:

#### a. Group I-Hardenable Chromium Steels (Martensitic and Magnetic)

These steels respond to heat treatment in a manner similar to most lower alloy steels and will, by suitable thermal treatment, develop a wide range of mechanical properties. The tempering range of 700° to 1000°F. should be avoided because it is conducive to temper brittleness.

#### b. Group II-Nonhardenable Chromium Steels (Ferritic and Magnetic)

These steels are essentially nonhardenable by heat treatment. They develop their maximum softness, ductility and corrosion resistance in the annealed condition.

c. Group III---Nonhardenable Chromium-Nickel and Chromium-Nickel-Manganese Steels (Austenitic and Nonmagnetic)

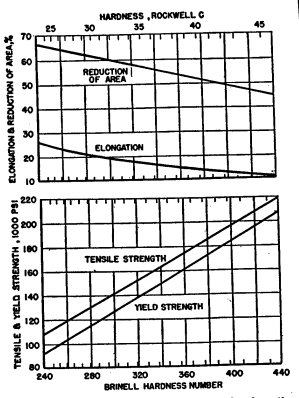
These steels are essentially nonmagnetic and cannot be hardened by heat treatment. Cold working develops a wide range of mechanical properties and some of these steels, in the severely cold worked condition, may become slightly magnetic. They are annealed by rapidly cooling (air or water) from high temperatures to develop maximum softness and ductility as well as corrosion resistance.

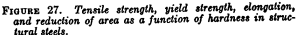
Recommended heat treatments for the steels listed in table 4 are given in table 8.

# 11. Properties and Uses of Steels

#### 11.1. Structural Steels

The strength properties of heat treated (quenched and tempered) structural carbon and alloy steels are closely related to their hardness and are surprisingly similar for a selected hardness provided that the steels originally were hardened throughout their cross sections. The relations of tensile strength, yield strength, elongation, and reduction of area to hardness are shown in figure 27. The tempering temperatures necessary to secure certain hardness levels in steels are affected by alloying elements, and two different steels may have to be tempered at two different temperatures to secure the same hardness. When this has been accomplished, however, the strength and ductility of the two steels will be quite similar. It must be emphasized that these relationships are valid only if the heat treated structures are tempered martensite. A steel that has been incompletely hardened (that is, guenched at too slow a rate to prevent the formation of some fine pearlite) may have, after tempering, a hardness and tensile strength equal to that of a completely hardened and tempered steel, but its yield strength and ductility will be inferior.





Valid only for steels originally hardened throughout and then tempered. Although drawn as lines, the relationship between these properties is not exact, and some deviations may be expected.

The modulus of elasticity of steel is the same as that of iron (about 29,000,000 lb/in.<sup>2</sup>). It is not affected by heat treatment or by the addition of alloying elements. Since stiffness, or the resistance to deformation under load, is a function of the modulus of elasticity, it follows that the stiffness of steel cannot be changed by heat treatment or by alloying elements, provided that the total stress is below the elastic limit of the steel in question. Either heat treatment or alloying elements can raise the elastic limit and thus apparently improve the stiffness in that higher allowable unit stresses may be imposed on the steel.

## a. Plain Carbon Structural Steels

The plain carbon steels are the least costly and may be used for a variety of purposes. The lower carbon grades (up to about 0.25% of carbon), frequently termed machinery steels, are used in the hot- and cold-worked conditions and for carburizing.Except when carburized, they are not very responsive to hardening by heat treatment because of the low carbon content. The medium carbon steels (about 0.30 to 0.60%) are the forging grades and are commonly used in the heat-treated condition. The higher carbon grades are the spring and tool steels.

Carbon steels are essentially shallow-hardening, usually requiring a water or brine quench, although very small sections may be successfully hardened in oil. Even with the most drastic quench, they cannot be hardened throughout even in moderate (about <sup>3</sup>/<sub>4</sub>-in.) cross sections. Large cross sections (about 4 in.) cannot be fully hardened even on the surface except by induction or flame hardening.

#### b. Alloy Structural Steels

From the viewpoint of heat treatment, the advantages accruing from the use of alloy steels are threefold: (a) Increased hardenability, (b) retention of fine-grained austenite, and (c) retardation of softening during tempering.

All of the alloying elements commonly used in structural alloy steels are effective to varying degrees in displacing the S-curve to the right as compared with plain carbon steels of similar carbon contents. Consequently, they can be hardened throughout in larger sections and, generally, by means of a milder quenching medium, such as oil. Because of the slower rates of quenching that can be employed in hardening these steels, they are less likely to distort or crack during hardening than the plain carbon steels.

The hardenability of alloy steels varies widely, depending on the composition. Certain elements and combinations of elements are very effective in increasing hardenability. Of the five alloying elements most generally used, and within the amounts normally present in the structural steels, nickel enhances hardenability but mildly, vanadium, chromium, manganese, and molybdenum moderately to strongly, depending upon the amount of alloy dissolved in the austenite at the hardening temperature. Combinations of chromium, nickel, and molybdenum, such as are present in the 4300 series (table 1), result in very deep hardening steels.

The alloy elements that form stable carbides, principally vanadium and molybdenum (and chromium to a lesser degree), are very effective in inhibiting grain growth in austenite. These steels may be overheated to a considerable degree during heating for hardening without suffering grain growth. This is desirable since martensite formed from fine-grained austenite is tougher than that formed from coarse-grained austenite.

Alloying elements, particularly those that form the more stable carbides, tend to retard the softening effects of tempering so that alloy steels must frequently be tempered at higher temperatures than carbon steels in order to secure the same hardness. This is beneficial in conferring increased toughness, i.e., resistance to shock. It is pertinent to note that manganese and nickel lower the annealing and hardening temperatures, and that chromium, molybdenum, and vanadium raise these temperatures. Combinations of these alloys may therefore, raise, lower or have no effect on the heat-treating temperatures, depending on the combinations of elements and their amounts.

The low carbon (up to about 0.25% of carbon) alloy steels are used mainly as carburizing steels. The medium carbon (about 0.30 to 0.60% of carbon) alloy steels find wide use as stressed members in an almost infinite variety of structural parts. Although some of these alloy steels containing high carbon are used for certain special applications, the maximum amount of carbon present in the alloy structural steels in generally about 0.5 percent.

#### 11.2. Tool Steels

Practical experience has shown that in a majority of instances the choice of a tool steel is not limited to a single type or even to a particular family for a working solution to an individual tooling problem. It is desirable to select the steel that will give the most economical overall performance; the tool life obtained with each steel under consideration should be judged by weighing such factors as expected productivity, ease of fabrication, and cost.

Hardness, strength, toughness, wear resistance, and resistance to softening by heat are prime factors that must be considered in selecting tool steel for general applications. Many other factors must be considered in individual applications; they include permissible distortion in hardening, permissible decarburization, hardenability, resistance to heat checking, machinability, grindability, and heat treating requirements.

The straight carbon tool steels can be used for a variety of purposes, depending upon the carbon content and heat treatment. The lower carbon ranges are used for tools where toughness and resistance to shock are of primary importance; these steels are usually tempered at temperatures of 500 to 700 °F or even higher. The higher carbon ranges are used where the main requirements are hardness, resistance to abrasion, or ability to hold a keen edge; these steels are usually tempered at temperatures of 300 to 500 °F.

An addition of 0.20 to 0.50 percent of chromium or of vanadium is frequently made to the carbon tool steels. The chromium-bearing carbon steels have a greater depth of hardening and are slightly more wear-resistant than the chromium-free steels of the same carbon content. Vanadium decreases slightly the depth of hardening but increases toughness. (This may appear contradictory to previous state-

ments that alloying elements increase hardenability. In the case of the carbon-vanadium tool steels, the normal hardening temperatures are too low to allow the solution of vanadium carbide in the austenite. The presence of undissolved carbides decreases hardenability).

The cutting ability, or the ability to hold an edge, is closely related to the hardness. For most purposes, therefore, cutting tools are used in a highly hardened condition. Excessive heat, whether caused by heavy cutting or careless grinding, will "draw the temper" (that is, decrease the hardness) and ruin the tool, necessitating rehardening and retempering. Since hardened carbon steels soften rapidly under the influence of heat, the carbon steels cannot be used as cutting tools under conditions where an appreciable amount of heat is generated at the cutting edge. Their uses are limited to conditions entailing light cuts on relatively soft materials, such as brass, aluminum, and unhardened low carbon steels.

The ranges for the carbon content and some typical applications of carbon tool steels are as follows:

0.60 to 0.75 percent of carbon—Hot forming or heading dies for short runs, machinery parts, hammers (sledges and pinch bars), concrete breakers and rivet sets.

0.75 to 0.90 percent of carbon—Hot and cold sets, chisels, dies, shear blades, mining drill steel, smiths' tools, set hammers, swages, and flatteners.

0.90 to 1.10 percent of carbon—Hand chisels, small shear blades, large taps, granite drills, trimming dies, drills, cutters, slotting and milling tools, mill picks, circular cutters, threading dies, cold header dies, jewelers' cold striking dies, blanking, forming and drawing dies, and punches.

1.10 to 1.40 percent of carbon—Small cutters, small taps, drills, reamers, slotting and planing tools, wood-cutting tools, turning tools, and razors.

Suitable tempering temperatures for carbon tool steels are as follows:

300 to 375 °F—Lathe tools and milling cutters, scrapers, drawing mandrels, dies, bonecutting tools, engraving tools, gages, and threading dies.

375 to 500  $^{\circ}$ F—Hand taps and dies, hand reamers, drills, bits, cutting dies, pen knives, milling cutters, chasers, press dies for blanking and forming rock drills, dental and surgical instruments, hammer faces, wood-carving tools, shear blades, and hack saws.

500 to 700 °F—Bending dies, shear blades, chuck jaws, forging dies, tools for wood cutting, hammers and drop dies, axes, cold chisels, coppersmith tools, screwdrivers, molding and planing tools, hack saws, butcher knives, and saws.

The high-speed steels are intended for use under heavy cutting conditions where considerable heat is generated. These steels are very complex, containing large amounts of alloying elements. The high-speed steels, which harden from a very high temperature (2,150 to 2,400 °F), develop what is termed "secondary hardness" after tempering at about 1,050 to 1,100 °F and maintain their cutting edge at considerably higher temperatures than do carbon tool steels.

Dies, depending on their use, can be made of a variety of steels. When intended for use at low temperatures, even carbon steel may be satisfactory sometimes. For use at elevated temperatures, certain minimum amounts of alloying elements are necessary and for the higher ranges of hot-working temperatures, steels of the high-speed steel type are frequently used.

The manufacture of dies usually involves such considerable expense in machining that cost of material and heat treatment form but a relatively small proportion of the total. It is imperative, therefore, that die steels be selected carefully and properly heat treated. The so-called "nondeforming" tool steels (type O). are favorites for dies that are not required to operate at elevated temperatures. These steels usually contain about 0.9 percent of carbon and 1.2 percent to 1.6 percent of manganese. With the lower manganese content they also contain about 0.5 percent each of chromium and tungsten. The expansion of these steels during hardening is much less than is experienced with carbon steels, and the shrinkage that occurs during the initial stage of tempering is almost sufficient to return the steel to its original size when it has been tempered at the proper temperature.

The high carbon-high chromium steels (type D) are frequently used for dies. This type contains about 12 percent of chromium and 1.5 or 2.2 percent of carbon, the lower carbon steel being air hardening. These steels also are non-deforming, are more resistant to wear, and may be used at higher temperatures than the manganese nondeforming steels. They have the disadvantage of requiring higher hardening temperatures (about 1,800 °F).

#### 11.3. Stainless and Heat Resisting Steels

Steels for high-temperature service must resist scaling and have high creep strength, that is, resistance to deformation under prolonged stress at elevated temperatures. Resistance to scaling is aided by the presence of chromium, aluminum, or silicon, and heatresistant steels invariably contain one or more of these elements. Resistance to creep is aided by elements that form stable carbides, such as tungsten, vanadium, molybdenum, and chromium.

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The stainless steels, often called corrosionresistant steels, contain appreciable amounts of chromium (12% or more), either with or without other elements, the most important of which is nickel. The heat-resistant and corrosion-resistant steels all belong to the same family, the terminology frequently being dictated by the temperature of use. By common consent, this dividing temperature is taken at about 1200 °F.

The three general groups of stainless steels have markedly different characteristics, as follows:

#### a. Group I-Hardenable Chromium Steels (Martensitic and Magnetic)

The significant characteristic of these is that the steels respond to heat treatment in a manner similar to most lower alloy steels. The steels possess a pearlitic structure on slow cooling and a martensitic (hardened) structure on rapid cooling from within the austenitic field. The chromium content varies from about 12 to 18 percent and the carbon content from less than 0.1 to over 1.0 percent; the higher chromium content demands the higher carbon.

The mechanical properties of this group of steels are markedly improved by heat treatment and heat treatment is essential to realize their corrosion-resistant properties. The only exceptions are the low carbon, low chromium steels (types 403, 410, 416) which are corrosion resistant in the annealed condition, although heat treatment decidedly improves mechanical properties.

Finishing of the martensitic steels has an important effect on corrosion resistance, since maximum corrosion resistance is obtained only with highly finished surfaces. Grinding should be done carefully, using a sharp wheel, light cuts, and plenty of coolant. Grinding burrs and embedded scale serve as focal points for corrosion.

As a group, the martensitic steels resist many types of corrosive environments, including the atmosphere, fresh water, mine water, steam, food acids, carbonic acid, crude oil, gasoline, blood, perspiration, ammonia, and sterilizing solutions.

Type 410 is a general purpose steel suitable for numerous applications where severe corrosion is not a problem. Types 416 and 416L are free-machining varieties of type 410. The higher hardness of type 420 makes it suitable for such uses as cutlery, surgical instruments, valves, ball bearings, magnets, etc. Type 420F is its free-machining counterpart. Where still greater hardness is required, type 440A, B, or C may be used; the last is the hardest of all the stainless steels. Types 420 and 440 should be used only in the hardened and stress-relieved conditions. Type 431 is a structural steel capable of being used in the strength range of 200,000 psi.

#### b. Group II-Nonhardenable Chromium Steels (Ferritic and Magnetic)

The distinguishing characteristic of this group is that these steels are ferritic at all temperatures and, therefore, incapable of being hardened by heat treatment. They can, however, be strengthened by cold working.

The ferritic steels develop coarse grained structures when subjected to temperatures above 1650 °F for varying lengths of time. This grain growth, frequently objectionable because of concomitant brittleness, cannot be eliminated by heat treatment alone. It can be corrected to some extent by cold working, followed by annealing.

The ferritic steels have lower strength at elevated temperatures than the martensitic steels, but resistance to scaling and corrosion is generally better. As a class, the ferritic stainless steels resist corrosion from food products, organic acids, salt water, nitric acid, many fused salts and molten nonferrous metals.

Type 430 finds extensive use in automobile trim, chemical equipment, food containers, and furnace parts. If ease of machining is important, type 430F or 430F Se may be substituted. Where greater resistance to corrosion or oxidation is required, type 446 may be used.

c. Group III-Nonhardenable Chromium-Nickel and Chromium-Nickel-Manganese Steels (Austenitic and Nonmagnetic)

The steels in this group are austenitic at room temperature and, hence, both nonhardenable by heat treatment and nonmagnetic. They can, however, be strengthened appreciably by cold work. Depending upon the amount of cold work, the tensile strength varies from about 80,000 psi for fully annealed material to as high as 300,000 psi for severely work hardened steel; the highest strengths can be secured only with small cross sections. A stress-relieving treatment of about 1 hr at 750 to 800 °F after severe cold working improves elastic properties without any adverse effect upon ductility. Cold work causes partial transformation of austenite to ferrite, with consequent appearance of magnetism, in some of these steels.

The austenitic steels have excellent coldforming properties and are extremely tough and ductile, even at low temperatures. Their toughness makes machining of all but the freemachining varieties difficult.

As a group, the austenitic steels are the most corrosion resistant of all the stainless steels. They have excellent resistance to acetic, nitric, and citric acids (liquid), foodstuffs, sterilizing

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solutions, most of the organic chemicals and dyestuffs, and a wide variety of inorganic chemicals. Typical applications are for outdoor trim, kitchen equipment, dairy utensils, soda fountains, and light weight transportation such as aircraft.

This group of steels should not be discussed without some mention of intergranular embrittlement. If maintained in the temperature range of about 800 to 1,400 °F, or if slowly cooled through this range after annealing, some of these steels precipitate carbides at the grain boundaries. This is not deleterious unless the steel is either simultaneously or subsequently exposed to acidified corrosive conditions, causing intergranular corrosion. When this occurs, the steels lose their metallic ring and become quite brittle.

Type 302, with its higher carbon content, is more prone to carbide precipitation than is 304, which in turn is more susceptible than 304L.

Prolonged exposure of these steels at temperatures within the range of 1000 to near 1600 °F may also promote the formation of a sigma phase constituent, resulting in increased hardness and decreased ductility, notch toughness and corrosion resistance.

Type 301, because of its lower alloy content, is quite susceptible to work hardening and is used largely in the cold rolled or cold drawn form of sheet, strip, and wire. Types 201 and 202 are similar to types 301 and 302 in properties. The former were developed primarily to conserve nickel.

Types 305, 308, 309, and 310 contain progressively higher amounts of chromium and nickel and are, therefore, superior to the lower alloy steels in corrosion and heat resistance.

Types 316 and 317 contain molybdenum. The incorporation of this element increases resistance to corrosion, particularly of the "pitting" type.

Types 321 and 347 were developed specifically to resist intergranular corrosion. This property of the titanium-bearing steel (type 321) is generally enhanced by a stabilizing heat treatment. This treatment is not particularly necessary for the niobium-bearing steels (types 347 and 348).

#### d. Precipitation-Hardenable Stainless Steels

The precipitation-hardenable stainless steels constitute a group of proprietary steels for which standard specifications have not been published by AISI. They may be grouped into three types; martensitic, semi-austenitic and austenitic. These steels are of increasing commercial importance because they can be strengthened by heat treatment after fabrication without resorting to very high temperatures or rapid quenches. They possess prop-

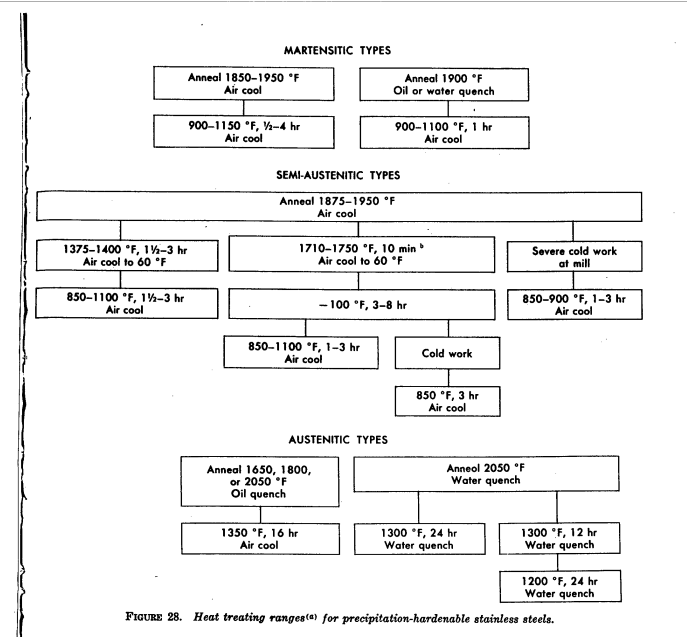
erties that make them particularly useful for structural parts and jet-engine components in high-performance aircraft as well as many non-military applications. Desirable characteristics include ease in hot working, machining, forming, and joining. High strength can be obtained by relatively-low elevated-temperature aging treatments or by refrigeration plus aging treatments. Ultrahigh strengths can be produced by a combination of cold working and aging. These steels usually possess good properties at ambient and moderately elevated temperatures (temperatures below those of the aging treatments) along with good corrosion and oxidation resistance. The austenitic types also retain very good impact properties at subzero temperatures.

Martensitic Types—The martensitic types are austenitic at annealing temperatures but transform to martensite of fairly low hardness on cooling to room temperature, either by air cooling, or by oil or water quenching. Elements contributing to the precipitation-hardening phases remain in supersaturated solid-solution in the steels on cooling. Precipitation occurs on the subsequent aging of the steels within the temperature range of 800 to 1100 °F (Figure 28). The martensitic matrix is also tempered by the aging treatment. The cooling method, aging temperatures and times at temperature depend upon the specific composition of the steel. Manufacturer's literature should be consulted for specific recommendations.

The ductility and impact strength of these hardened martensitic steels are relatively low compared to those of either the semi-austenitic or austenitic types. Increasing the aging temperature raises the ductility and impact strength with an accompanying loss in the yield and tensile strengths. These steels, in the annealed condition, lack formability as compared with that of the semi-austenitic or austenitic types of precipitation-hardenable stainless steels.

Semi-Austenitic Types—The semi-austenitic steels are austenitic as annealed within the temperature range of 1850 to 1950 °F and air cooled to room temperature (large sections may be water quenched). They are normally supplied by the mill in the annealed condition or in an annealed plus severe cold-worked condition. By careful control of composition and annealing treatment, an austenite can be produced which has a stability intermediate between that of the martensitic and austenitic types of precipitation-hardenable stainless steels. These steels are readily amenable to forming operations after the annealing treatment.

Some representative heat treatments for these steels are shown in figure 28. The series of treatments (temperatures and times at tem-



(a) Ranges of temperatures, heating periods and cooling rates (quenching mediums) shown include those for several commercial steels. Heat treatment procedures and values depend upon the composition of the specific steel selected. These steels are generally provided in the annealed (solution-treatd) condition by the mills. in the annealed (solution-treated) condition by the mills.

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perature) to be followed depend upon the specific composition of the steel and the desired final properties. After the forming operations, these steels are usually given a destabilizing treatment at temperature ranges of 1375 to 1400 °F or 1710 to 1750 °F in which some of the carbon is rejected from the solid solution as a carbide. This facilitates the transformation of the austenite to martensite during the cooling to 60 °F or -100 °F and results in a strengthening of the matrix. However, the final high strengths of these steels are obtained by the subsequent aging treatments at temperatures ranging form 850 to 1100 °F. The aging treatments temper the martensite and

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increase the strength of the steels by precipitation of solute elements from solid solution as hardening phases (usually nickel-aluminum compounds). Cold working of the steel following the transformation of the austenite to martensite and prior to the aging treatment produces very high strengths.

If these semi-austenitic steels are severely cold-worked (usually at the mill) following the annealing treatment most of the austenite is transformed to martensite by the cold working. Then no destabilizing treatment is needed prior to the aging treatment which produces very high strengths. However, the retained ductility is usually very low.

These initially semi-austenitic steels through the hardening and aging treatments develop the highest room-temperature tensile properties (tensile strengths up to nearly 300,000 psi) of the three types of precipitation-hardenable stainless steels. Manufacturer's literature should be consulted for heat treatment procedures as they depend very strongly on the specific composition of the steel and its intended use.

Austenitic Types-The austenitic steels are austenitic in both the annealed and hardened conditions. They are normally supplied by the mill in the annealed condition. Annealing temperatures (Figure 28) vary from 1650 to 2050 °F, depending upon the composition of the steel. The steels are cooled rapidly from the annealing temperatures either by oil or water quenching. These steels are subjected to the forming or cold working operations prior to hardening. The hardening treatment (Figure 28) consists of aging within the tem-perature range of 1200 to 1400 °F which precipitates carbides and other hardening phases. Cold working of these steels prior to aging increases the strength with the properties depending upon both the degree of cold-working and the aging treatment. As the heat treatment procedure for these steels depends upon the composition of the steel and its intended use, the manufacturer's literature should be consulted for details of chemical composition, recommended annealing and aging treatments, and typical properties and uses.

In general the austenitic steels have lower room-temperature properties than either the martensitic or semi-austenitic types. However, they retain very good tensile and impact properties at subzero temperatures. Moreover, they retain be'ter tensile, creep and rupture properties at temperatures up to their aging temperatures than the other two types of precipitation-hardenable stainless steels.

#### 11.4. Nickel Maraging Steels

A series of high-strength iron-base alloys of very low carbon content (0.03%) with yield strengths of 250,000 psi or greater in combination with excellent fracture toughness have been developed in recent years. These are the nickel maraging steels which do not require rapid quenching, as the hardening and strengthening are obtained by an iron-nickel martensite transformation and an age-hardening reaction. The iron-nickel martensite is only moderately hard (approximately Rockwell C 25) and very tough compared to that of untempered carbon steel martensites. The transformation of austenite to martensite is not sensitive to cooling rates. Thus rapid quenching is not required and section-size effects are small. No significant tempering occurs upon

the reheating of the iron-nickel martensite. Moreover, no appreciable reversion of martensite to austenite occurs during the reheating of the steel for the aging reaction (precipitation of intermetallic compounds\* in the presence of suitable alloying elements).

The maraging steels are proprietary steels for which standard specifications have not been published by AISI. Four grades are commercially available: 25 Ni (25% Ni, 1.3-1.6% Ti); 20 Ni (20% Ni, 1.3-1.6% Ti); 18 Ni (17-19%Ni, 7-9.5\% Co, 3-5% Mo, and 0.2-0.8% Ti); and 12 Ni (9-15% Ni, 3-5% Cr, 2-4% Mo, and 0.08-0.55\% Ti). All of these steels contain appreciable amounts of aluminum. The 25 Ni and 20 Ni grades have a niobium content of 0.30 to 0.50\%.

The maraging steels can be heat treated in conventional furnaces without protective atmospheres. They can be formed or machined to final dimensions in the annealed condition and will retain close tolerances during the aging treatment. Oxidation during aging is usually insignificant. Heat treating ranges are showr in figure 29. As the heat treatment proce dures depend upon the specific composition of the steel, the manufacturer's literature shouk be consulted for the chemical composition and recommended heat treatment of the selected steel.

#### a. 25 Ni Maraging Steel

The 25 Ni steels remain austenitic upon ai cooling to room temperature after annealing (solution-anneal) at 1500 °F. The steels i this condition are relatively soft and readily fabricated. As indicated in figure 29, two prc cedures are available for conditioning the steel for transformation to martensite. The steel may be given an aging treatment at 1300 °] which reduces the stability of the austenit presumably by precipitation of nickel-titaniur compounds in the austenite. This raises th martensite transformation temperature so the the austenite will transform to martensite o cooling to room temperature. In the othe procedure, the martensitic transformation ca be induced by cold working the annealed stee to a reduction of 25% or more. After either treatment, cooling at -100 °F for sever hours secures complete transformation. Fu strength properties (250,000-270,000 psi yie strength) are then obtained by the agin treatment at 900 °F for 1 to 3 hours.

#### b. 20 Ni Maraging Steel

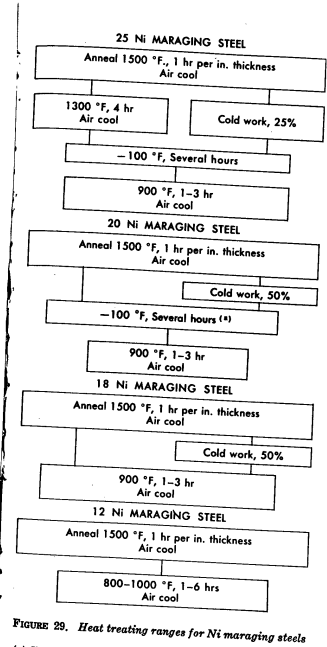
The annealed 20 Ni steels have a martensit transformation temperature above room ter perature and hence they transform to marter

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<sup>\*</sup> Currently, there is a question as to whether or not order occurs during the aging treatment and contributes to the streng ening of the steel.



(a) Usually recommended, but may be omitted.

ite upon cooling to room temperature from the annealing or hot working temperatures (figure 29). Refrigeration at -100 °F for several hours is usually recommended for assurance of complete transformation prior to the aging treatment at 900 °F for 1 to 3 hrs. For enhanced strength (yield strength up to 260,000 psi) the steels may be cold worked following the annealing and prior to refrigeration at -100 °F and subsequent aging.

# c. 18 Ni Maraging Steel

The heat treatment procedures for the 18 Ni steels are the same as those for the 20 Ni steels except that refrigeration of the steels is not necessary; the complete transformation of the austenite to martensite occurs during air cooling of the steels from the hot working or annealing temperatures to room temperature. Full strength properties are obtained by the aging treatment at 900 °F. For the highest strength properties (yield strength of 300,000 to 310,000 psi) the steels are cold worked to reductions of 50% or more following the annealing and prior to the aging treatment.

# d. 12 Ni Maraging Steel

The 12 Ni steels combine intermediate strengths (150,000-215,000 psi yield strength) with good ductility and high toughness as measured by Charpy V-notch impact resistance at room temperature. These steels usually retain good impact resistance at subzero temperatures down to -100 °F. The heat treating procedures are indicated in figure 29. They are very similar to those for the higher strength 18 Ni steels.

In preparing this monograph, the authors have drawn liberally from the field of common knowledge and published information, especially from the Metals Handbook, published by the American Society for Metals, and the Steel Product Manuals of the American Iron and Steel Institute. Grateful acknowledgment is due to all metallurgists, both known and unknown, whose work and study have transformed the art of heat treating into a science. The various photomicrographs used were prepared by Carolyn R. Irish, a former member of the staff of the National Bureau of Standards

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