2019 Fall

"Advanced Physical Metallurgy" - Non-equilibrium Solidification -

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Glass Formation results when

Liquids are cooled to below $T_m(T_L)$ sufficiently fast to avoid crystallization.

Nucleation of crystalline seeds are avoided

Growth of Nuclei into crystallites (crystals) is avoided

Liquid is "frustrated" by internal structure that hinders both events

"Glass Formation"

Critical cooling rate is inversely proportional to the diameter of ingot.



Critical Cooling Rates for Various Liquids

Table 3-5. Examples of Critical Cooling Rates (°C/s) for Glass Formation

Material	Homogeneous nucleation	Heterogeneous nucleation contact angle (deg)		
		100	60	40
SiO_2 glass ^a	9×10^{-6}	10^{-5}	8×10^{-3}	2×10^{-1}
GeO_2 glass ^a	3×10^{-3}	3×10^{3}	1	20
Na ₂ O·2SiO ₂ glass ^a	\bullet 6 × 10 ⁻³	8×10^{-3}	10	$3 \times 10^{+2}$
Salol	10			
Water	107			
Ag	10 ¹⁰			
Typical metal ^a	9×10^8	9×10^9	10 ¹⁰	5×10^{10}

^a After P. I. K. Onorato and D. R. Uhlmann, J. Non-Cryst. Sol., 22(2), 367–378 (1976).

Q4: Overall Transformation Kinetics-TTT diagram

"Johnson-Mehl-Avrami Equation"

5.4 Overall Transformation Kinetics – TTT Diagram

If isothermal transformation,

The fraction of Transformation as a function of Time and Temperature

 $\rightarrow f(t,T)$

 $\operatorname{Plot} f$ vs log t.

- isothermal transformation
- $f \sim$ volume fraction of β at any time; $0 \sim 1$

Plot the fraction of transformation (1%, 99%) in T-log t coordinate.



Fig. 5.23 The percentage transformation versus time for different transformation temperatures.

Influence factors for *f*(t,T): **nucleation rate, growth rate, density and distribution of nucleation sites, impingement of adjacent cells**

<u>Three Transformation Types</u>







(a) Beginning





(b) Nucleation sites



(c) Cellular transformation

Fig. 5.24 (a) Nucleation at a constant rate during the whole transformation.(b) Site saturation – all nucleation occurs at the beginning of transformation.(c) A cellular transformation.

(a) continuous nucleation

Metastable α phase with many nucleation sites by quenching to T_t

- \rightarrow f depends on the nucleation rate and the growth rate.
- (b) all nuclei present at t = 0 $\rightarrow f$ depends on the number of nucleation sites and the growth rate.
- (c) All of the parent phase is consumed by the transformation product.

Transformation terminate by the impingement of adjacent cells growing with a constant velocity.

→ pearlite, cellular ppt, massive transformation, recrystallization

Transformation Kinetics

Avrami proposed that for a three-dimensional nucleation and growth process kinetic law

$$f=1-\exp(-kt^n)$$
 Johnson-Mehl-Avrami equation

f : volume fraction transformed

 $=\frac{\text{Volume of new phase}}{\text{Volume of specimen}}$

> Assumption :

- \checkmark reaction produces by nucleation and growth
- \checkmark nucleation occurs randomly throughout specimen
- \checkmark reaction product grows rapidly until impingement

Constant Nucleation Rate Conditions

Nucleation rate (/) is constant. (*t*) Growth rate (v) is constant. No compositional change t τ $df_{e} = \frac{\begin{pmatrix} \text{Vol. of one particle nucleated} \\ \text{during } d\tau \text{ measured at time t} \end{pmatrix} \times \begin{pmatrix} \text{number of nuclei} \\ \text{formed during } d\tau \end{pmatrix}}{\text{Volume of specimen}}$ $df_e = \frac{\frac{4}{3}\pi [v(t-\tau)]^3 \times (IV_0 d\tau)}{V_0}$ $f_e(t) = \int_0^t I \cdot \frac{4}{3}\pi [v(t-\tau)]^3 d\tau$ $V = \frac{4}{3}\pi r^{3} = \frac{4}{3}\pi (vt)^{3}$ $V' = \frac{4}{2} \pi v^3 (t-\tau)^3$ $= I \cdot \frac{4}{3} \pi v^3 \left[-\frac{1}{4} (t-\tau)^4 \right]_0^t = \left[\frac{1}{3} \pi I v^3 t^4 \right]_0^t - \frac{1}{3} dv^3 t^4 - \frac{1}{3$

As time passes the β cells will eventually impinge on one another and the rate of transformation will decrease again.

Constant Nucleation Rate Conditions

consider impingement + repeated nucleation effects



2.5.2 Kinetics of Glass Formation

A. Homogeneous Nucleation rate, *I* (by David Turnbull)

$$I = \frac{k_{\rm n}}{\eta(T)} \exp\left[-\frac{b\alpha^{3}\beta}{T_{\rm r}(\Delta T_{\rm r})^{2}}\right]$$

where

b is a shape factor (= $16\pi/3$ for a spherical nucleus)

 $k_{\rm n}$ is a kinetic constant

 $\eta(T)$ is the shear viscosity of the liquid at temperature T

 $T_{\rm r}$ is the reduced temperature ($T_{\rm r} = T/T_{\rm l}$)

 $\Delta T_{\rm r}$ is the reduced supercooling ($\Delta T_{\rm r} = 1 - T_{\rm r}$)

 α and β are dimensionless parameters related, respectively, to <u>the liquid</u>/ solid interfacial energy (σ) and to the molar entropy of fusion, $\Delta S_{\rm f}$

Thus, $\alpha = \frac{\left(N_{\rm A}\overline{V}^2\right)^{1/3}\sigma}{\Delta H_{\rm f}} \qquad \beta = \frac{\Delta S_{\rm f}}{R}$

where

 $N_{\rm A}$ is Avogadro's number \overline{V} is the molar volume of the crystal R is the universal gas constant

(2.4)

A. Homogeneous Nucleation rate, *I* (by David Turnbull)

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$$\alpha = \frac{\left(N_{\rm A}\overline{V}^2\right)^{1/3}\sigma}{\Delta H_{\rm f}} \qquad \beta = \frac{\Delta S_{\rm f}}{R}$$

- 1) $\eta \uparrow (\text{dense random packed structure}) \rightarrow I \downarrow$
- 2) For given T and η , $\alpha^{3}\beta$ \uparrow (σ solid interfacial E & $\Delta S_{f}\uparrow/\Delta H_{f}\downarrow$) $\rightarrow I\downarrow$
- 3) $\eta \sim T_{rg} (=T_g/T_l) \& \alpha^3 \beta \sim$ thermal stability of supercooled liquid * For metallic melt : $\alpha \beta^{1/3} \sim 0.5$
 - * if $\alpha\beta^{1/3}$ > 0.9, impossible to crystallization by homogeneous nucleation under any cooling condition
 - * if $\alpha\beta^{1/3} \leq 0.25$, difficult to prevent crystallization

B. Growth rate of a crystal from an undercooled liquid, U

$$U = \frac{10^2 f}{\eta} \left[1 - \exp\left(-\frac{\Delta T_r \Delta H_f}{RT}\right) \right]$$
(2.7)

where *f* represents the fraction of sites at the crystal surfaces where atomic attachment can occur (=1 for close-packed crystals and 0.2 ΔT_r for faceted crystals). Here also we can see that *U* decreases as η increases, and will thus contribute to increased glass formability.

- 1) $\eta \uparrow$ (dense random packed structure) $\rightarrow U \downarrow$
- 2) For given T , *I* & $U \sim \eta \rightarrow T_{rg}$ or α , $\beta \uparrow \rightarrow GFA \uparrow$
- 3) $f \downarrow$ through atomic rearrangement like local ordering or segregation $\rightarrow U \downarrow$
 - * metallic melt: $\alpha\beta^{1/3} \sim 0.5$ / $T_{rg} > 2/3 \sim$ high GFA
 - * Pure metal: $R_c \sim 10^{10-12}$ K/s, but if T_{rg} = 0.5 , $R_c \sim 10^6$ K/s

Based on the treatment of Uhlmann [25], Davies [26] combined the values of I and U (calculated using Equations 2.4 and 2.7, respectively) with the Johnson–Mehl–Avrami treatment of transformation kinetics, and calculated the fraction of transformed phase x in time t, for small x, as

C.
$$x = \frac{1}{3}\pi I U^3 t^4$$
 (2.8)

Substituting the values of *I* and *U* in Equation 2.8, the time needed to achieve a small fraction of crystals from the melt was calculated as

$$t \approx \frac{9.3 \eta a_o^2 x}{kT f^3 \overline{N}_v} \left[\frac{\exp\left(\frac{1.07}{\Delta T_r^2 T_r^3}\right)}{\left\{1 - \exp\left(-\frac{\Delta H_f \Delta T_r}{RT}\right)\right\}^3} \right]^{1/4}$$
(2.9)

where

 $a_{\rm o}$ is the mean atomic diameter

 \overline{N}_{v} is the average volume concentration of atoms, and all the other parameters have the same meaning, as described earlier

A time–temperature–transformation (T-T-T) curve was then computed by calculating the time, *t*, as a function of T_r , to transform to a barely detectable fraction of crystal, which was arbitrarily taken to be $x = 10^{-6}$.

Nucleation and Growth Rates – Poor Glass Formers



Nucleation and Growth Rates – Good Glass Formers



TTT curves and the critical cooling rate, R_c





FIGURE 2.3

Time-temperature-transformation (T-T-T) curves (solid lines) and the corresponding continuous cooling transformation curves (dashed lines) for the formation of a small volume fraction for pure metal Ni, and Au₇₈Ge₁₄Si₈, Pd₈₂Si₁₈, and Pd₇₈Cu₆Si₁₆ alloys.



FIGURE 10.11 A more complete TTT diagram for eutectoid steel than was given in Figure 10.7. The various stages of the time-independent (or diffusionless) martensitic transformation are shown as horizontal lines. M_s represents the start, M_{50} represents 50% transformation, and M_{90} represents 90% transformation. One hundred percent transformation to martensite is not complete until a final temperature (M_f) of $-46^{\circ}C$.

* Continuous Cooling Transformation diagrams



FIGURE 10.14 A continuous cooling transformation (CCT) diagram is shown superimposed on the isothermal transformation diagram of Figure 10.11. The general effect of continuous cooling is to shift the transformation curves downward and toward the right. (After Atlas of Isothermal Transformation and Cooling Transformation Diagrams, American Society for Metals, Metals Park, OH, 1977.)

Q5: Measurement of TTT diagram

Measurement of TTT Diagram during Heating by DSC/DTA





Study of hidden kinetics via fast heating



- Heating rate : 4*10⁴ K/s
- Cooling rate : 4*10³ K/s
- Temperature range:
 - -90°C 1000°C





UFS -90 – 450 °C Limited heating rate ~ 3*10⁴ K/s

Stable data regardless of sample weight

UFH -90 – 1000 °C

Sample weight less than 100ng is recommended

Extreme experimental condition \rightarrow hidden kinetics of materials



Measurement of TTT Diagram during Heating by DSC/DTA



Calculation of Time-Temperature-Transformation Diagram

Nucleation Rate

I = nvexp(-NW*/RT)exp(- $\Delta E_D/RT$)

W*: thermodynamic energy barrier to form nuclei

 $W^* = 16\pi\sigma^3/3(\Delta G_{cryst}(T))^2$

I = n ν exp(-16πσ³/3($\Delta G_{cryst}(T)$)²/RT) exp(- ΔE_D /RT)

Growth rate

 $\mu = avexp(-\Delta E_D/RT) \times (1 - exp(\Delta G_{cryst})/RT)$

D: diffusion coefficient E_D : diffusion energy barrier to form nuclei $D(T) = a^2 vexp(-\Delta E_D/RT) = fRT/3N\pi a^2\eta(T))$ Stokes-Einstein relation between D and η $\mu(T) = (fRT/3N\pi a^2\eta(T)) (1 - exp(\Delta G_{cryst})/RT)$



Calculation of Time-Temperature-Transformation Diagram

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D : diffusion coefficient E_D : diffusion energy barrier to form nuclei D(T) = a²vexp(- ΔE_D/RT) = fRT/3Nπa²η(T))

Stokes-Einstein relation between D and η

 $\mu(T) = (fRT/3N\pi a^2\eta(T)) (1 - exp(\Delta G_{cryst})/RT)$





TTT Diagrams of Pd₄₀Cu₃₀Ni₁₀P₂₀

Measurement of TTT Diagram by Drop Tube Technique

Solidification of containerless undercooled Melts, edited by Dieter M. Herlach and Douglas M. Matson, 2012, p.1-7



Drop tube technique

- rapid cooling of small particles by dispersion of the melt
- reduction of heterogeneous nucleation by containerless processing



TPhasegraxusrofitheroptetssopAlgglossipvallogd in the solid tissation of om de choopled disoptets of Alg8Mn12

Electrostatic Levitation in KRISS



Electrostatic Levitation in KRISS



Measurement of TTT Diagram by ESL Technique



TTT Diagram of Pd_{42.5}Ni_{7.5}Cu₃₀P₂₀



TTT Diagram of Pd₄₀Cu₃₀Ni₁₀P₂₀



2.6 Methods to Synthesize Metallic Glasses

2.6.1 Vapor-state Processes: expensive & slow, electronic & magnetic applications Thermal Evaporation/ Sputtering/ Vapor Chemical Deposition



2.6.2 Liquid-state Processes : Rapid Solidification Process (RSP) 10⁵⁻⁶ K/s most ideal way to obtain metallic glasses, especially the bulk foam

Splat Quenching/ Melt-spinning/ Electro Deposition/ Gas Atomization

2.6.3 Solid-state Processes_Solid state diffusional amorphization

Mechanical Alloying & Milling/Hydrogen-induced Amorphization/ Multilayer Amorphization/ Pressure-induced Amorphization/ Amorphization by Irradiation/ Severe Plastic Deformation_Intense deformation at low temperatures /Accumulative Roll Bonding (ARB process) 2.6 Methods to Synthesize Metallic Glasses

2.6.1 Vapor-state Processes: expensive & slow, electronic & magnetic applications

A. Thermal Evaporation

Surface coating

- Thermal evaporation is a common method for thin film deposition.
- The source material is evaporated from a source by heating in a vacuum.
- The evaporated material is then condensed on the substrate.





Heating : resistance heating (Joule effect)

First Amorphous Metals: evaporation method

Über nichtleitende Metallmodifikationen¹)

Von Johannes Kramer

(Mit 8 Figuren)

Das metallische Leitvermögen wird bekanntlich auf das Vorhandensein freibeweglicher Elektronen und damit auch ortsgebundener positiver Ionen zurückgeführt. Da nun ein nichtionisierter Metalldampf ein vollkommener Nichtleiter ist, so liegt die Vermutung nahe, daß es bei Kondensation eines solchen Dampfes gelingen müßte, nichtleitende Schichten zu erhalten, wenn Wechselwirkungen zwischen den regellos aufeinandergepackten Atomen vermieden werden könnten. Man hätte es dann mit einem Gebilde zu tun, das als völlig amorph anzusehen wäre und in seiner Konstitution am ehesten einem hochkomprimierten Gase entspräche.





Fig. 1. Zerstäubungsapparatur

J. Kramer Nonconducting modifications of metals. Ann. Physik (Berlin, Germany) 19, 37 (1934)

Sb metal \rightarrow Bi, Ga, and Sn-Bi alloys
What is Sputtering?

Surface coating

- B. Sputtering is a common method for thin film deposition.
- Process whereby atoms are ejected from a solid target material due to bombardment of the target by energetic particles.
- Ejected atoms are deposited on the substrate, form thin film.

Ref. http://en.wikipedia.org/wiki/Sputtering

Vacuum : evaporated particles can travel directly to the substrate



Radio Frequency Sputtering

Inert **RF** Power Gas Electrode Target å plasma A Substrate Electrode(Hot chuck) Vacuum Pump Neutral target atom lonized atom Electron

DC vs RF

	DC Sputter	RF Sputter
Deposition rate	faster	slower
Inert Gas Pressure	Higher	lower
Dep.material	Metal	Metal, Insulator, oxide
ETC		Improve plasma handling.

Controlling Temperature of Chuck Cooling rate of the thin film can be modulated. Some of materials can formed to amorphous state.



Surface coating

C. 📴 Principles of CVD

- Substrate is exposed to one or more volatile precursors.
- Precursors will react and/or decompose on substrate to produce the desire deposit.

 $3 \operatorname{SiH}_4 + 4 \operatorname{NH}_3 \rightarrow \operatorname{Si}_3\operatorname{N}_4 + 12 \operatorname{H}_2$

Surface coating

144600





Surface coating

- PECVD (Plasma Enhanced CVD)
- LPCVD (Low Pressure CVD)
- APCVD (Atmospheric Pressure CVD)
- UHVCVD (Ultra-high vacuum CVD)
- DLICVD (Direct liquid injection CVD)
- AACVD (Aerosol-assisted CVD)
- MPCVD (Microwave Plasma CVD)
- ALCVD (Atomic Layer CVD)

Laser CVD

Formation of Amorphous Thin Film Sur

Surface coating

Calculation of Diffusion Length



Formation of amorphous

- Amorphous forming ability is depends on diffusion length
- ✤ Diffusion length $\propto \sqrt{Dt}$
- Deposit process is commonly forming amorphous because of low temperature and rapid growth rate
- CVD process is progressed at little high temperature, so metal do not forming amorphous generally



Surface coating



2.6.2 Liquid-state Processes : Rapid Solidification Process (RSP) 10⁵⁻⁶ K/s most ideal way to obtain metallic glasses, especially the bulk variety



Glass formation : stabilizing the liquid phase

First metallic glass (Au₈₀Si₂₀) produced by splat quenching at Caltech by Pol Duwez in 1960.



2.6.2 Liquid-state Processes

B. Brief Definition of Melt Spinning

A jet of liquid metal is ejected from a nozzle and impinges on the surface of a rotating substrate, where a thin layer is formed from a melt puddle and rapidly solidifies as a continuous ribbon.

Several Types of Melt Spinning Method



• Melt-spinning method

Thin film



Surface coating



C. Basic Concept of Electrolytic Deposition

Electroplating is a plating process in which metal ions in a solution are moved by an electric field to coat an electrode. The process uses electrical current to reduce cations of a desired material from a solution and coat a conductive object with a thin layer of the material, such as a metal. Electroplating is primarily used for depositing a layer of material to bestow a desired property (e.g., abrasion and wear resistance, corrosion protection, lubricity, aesthetic qualities, etc.) to a surface that otherwise lacks that property. Another application uses electroplating to build up thickness on undersized parts. However, the limited size of the product formed in one experiment puts a limitation on the usefulness of this technique to produce large quantities of metallic glasses



2.6.2 Liquid-state Processes

C. Electro-deposition Methods It was in 1950, for the first time, that Brenner et al. [38] reported the formation of an amorphous phase in electro-deposited Ni–P alloys containing >10 at.% P. The amorphous nature of the deposit was inferred from the presence of only one broad diffuse peak in the XRD pattern. These alloys have a very high hardness and consequently these are used as wear- and corrosion-resistant coatings [39].

U. S. Department of Commerce National Bureau of Standards Research Paper RP2061 Volume 44, January 1950

Part of the Journal of Research of the National Bureau of Standards

Electrodeposition of Alloys of Phosphorus with Nickel or Cobalt

By Abner Brenner, Dwight E. Couch, and Eugenia Kellogg Williams

Alloys containing nickel or cobalt and as much as 15 percent of phosphorus have been electrodeposited from solutions containing phosphites. The alloys are hard and may be further hardened by heat-treating at 400° C. The high-phosphorus nickel alloy is more resistant to attack by hydrochloric acid than pure nickel deposits. The high-phosphorus alloys are bright as deposited, but their reflectivities are lower than those of buffed coatings of pure nickel.

Powder

D. Gas Atomization



A. Mechanical alloying/milling

: MA/ MM performed in a high-energy ball mill such as a shaker mill or planetary mill will also induce severe plastic deformation in metals. During milling, particles are fractured and cold welded together, resulting in large deformation.

Schematic Diagram of Mechanical Alloying Process



Increased Solid Solubility Limit Easy to Control the Microstructure Easy to Fabricate Non-Equilibrium and Nano Phase Homogeneous Distribution of Fine Strengthener Excellent Mechanical Properties



A. Mechanical alloying/milling

The milled powder particles experience heavy plastic <u>deformation</u> leading to the generation of a variety of crystal defects such as dislocations, grain boundaries, and stacking faults. These defects <u>raise the</u> <u>free energy of the crystalline</u> system to a level higher than that of a hypothetical amorphous phase, and consequently, the crystalline phase becomes destabilized and an amorphous phase forms [41,42].

The first report of the formation of an amorphous phase by mechanical milling was in a Y–Co intermetallic compound [43] and that by mechanical alloying in a Ni–Nb powder blend [44]. Subsequently, amorphous phases have been formed in a number of binary, ternary, and higher-order systems by this method. Unlike in RSP methods, the conditions for the formation of an amorphous phase by mechanical alloying/milling seem to be quite different. For example, amorphous phases are formed, not necessarily near eutectic compositions, and in a much wider composition range. The nature and transformation behavior of the amorphous phases formed by RSP.

A. Mechanical alloying/milling

Amorphization during Mechanical Alloying



Magnesium Technology Innovation Center

Mechanical Alloying/ Milling





B. Hydrogen-induced Amorphization

- Yeh et al. [45] reported the formation of an amorphous metal hydride Zr₃RhH_{5.5} by a reaction of hydrogen with a metastable crystalline Zr₃Rh compound at sufficiently low temperatures (< about 200°C). A similar amorphous hydride phase was also reported to form when the amorphous Zr₃Rh alloy obtained in a glassy state was also reacted with hydrogen.
- Aoki [46] has reported that hydrogen-induced amorphization is possible in many binary metal compounds.

C. Multilayer Amorphization

• An amorphous phase was found to form when thin metal films (10–50 nm in thickness) of La and Au were allowed to interdiffuse at relatively low temperatures (50°C–100°C). Similar to the case of hydrogen-induced amorphization, here also the large asymmetry in the diffusion coefficients of the two elements was found to be responsible for the formation of the amorphous phase [47]. Since then, a large number of cases where multilayer amorphization occurs have been reported.

Ex) cold rolling of thin foils of Ni and Zr, and then annealing them at a low temperature.

Solid-State Diffusional Amorphization

- Alternate layers of crystalline metallic films ('diffusion couples') interdiffuse under isothermal conditions, with the eventual amorphization of the entire multilayer.
- Couple: late transition metal with early transition metal ex. Au-La, Au-Zr, Au-Y, Cu-Zr, Cu-Er, Ni-Er, Ni-Ti, Ni-Hf, Fe-Zr, Co-Zr, Ni-Zr
- ✓ Low temperature annealing below the crystallization temperature (the nucleation or growth of crystalline phases cannot occur)

* Diffusion in multiple binary system



Solid-State Diffusional Amorphization

Schematic Diagram



- (A) Direct nucleation of the amorphous phase
- (B) Substantial mutual dissolution of the parent metals leading to unstable solid solutions

Two Main Attributes

A strong thermodynamic driving force

i.e. a large negative heat of mixing



Diffusional asymmetry

i.e. one element diffuses anomalously fast in the other, but not vice versa



D. Pressure-induced Amorphization

Alloys in the systems Cd–Sb, Zn–Sb, and Al–Ge were subjected to high pressures when they formed unstable crystalline phases and then decayed in a short time to amorphous phases [50]. An amorphous phase was also found to form in Cu–12–17 at.% Sn alloys when they were heated to high temperatures in a confined pressure, and turning off the power suddenly.

e. Amorphization by Irradiation

A number of intermetallics, for example, NiTi, Zr₃Al, Cu₄Ti₃, CuTi₂, FeTi, etc., have been amorphized by irradiation with high-energy electrons, heavy ions, or fission fragments [18]. It is only intermetallics, and not solid solutions, that have been amorphized this way. Several different criteria have been proposed for this purpose, which include low temperature, high doses, high dose rates, intermetallics with an extremely narrow homogeneity range, permanently ordered compounds (i.e., intermetallics which are ordered up to the melting temperature and do not show an order–disorder transformation), which also have high-ordering energies, destruction of long-range order by irradiation, etc. Fecht and Johnson [52] have analyzed the fundamental issues involved in the amorphization of intermetallics by irradiation methods.

Preparation of amorphous materials by irradiation Two kinds of irradiation effects

• Laser/electron irradiation:

Small area melting \rightarrow cooling \rightarrow amorphous layer/crystalline



Neutron and heavy ion irradiation:
Atomic displacement → recovery → amorphous phase

> Mechanism of irradiation amorphization:

- The kinetic knock-off of atoms from their lattice positions
- Very high temperature is expected in a very small volume during a short time (ps), which causes melting of material in a local small point.

> Two opposing radiation-induced processes can be operative:

- **Radiation-induced chemical disordering** of atoms on lattice sites will tend to promote amorphization;
- Radiation-induced defect migration will tend to restore ordering.
- ➡ The irradiation be <u>carried out at sufficiently low temperature</u> to suppress the defect migration.

Radiation damage: damage by collision between high energy particle and materials

• Basic defect by collision with high E particle: Vacancy & Interstitial

(Frenkel pair)



20keV Fe displacement cascade in bcc Fe at 600K

http://www.youtube.com/watch?v=0btHd_8JFV4

* A primary knock-on atom (PKA) displaces neighbouring atoms, resulting in <u>an atomic displacement</u> <u>cascade</u>, leading to formation of point defects and defect clusters of vacancies and interstitial atoms.

Seeger et al., Proc. Symp. Radiat. Damage Solids React. 1 (1962) 101-1056.

Radiation damage: damage by collision between high energy particle and materials



S. Zinkle, Defects, deformation and damage in structural materials, 2012

Sample Preparation : Surface Damage

Ion beam induced amorphous formation in Silicon (100)



Y.W. Kim, Metals and Materials International, 7, 499 (2001)

The schematic diagram of High Energy Electron beam accelerator





TT1000 prototype picture



The making method of amorphous materials by Irradiation



Electron beam irradiation is heating the mixture of powder.

✤ The temperature of substrate is so low that surface is super-cooling.



Surface : Amorphous alloy

Surface gloss, Hardness, corrosion resistance,

Substrate : Cu(or Ti, Fe) alloy

Ductility, limited size of amorphous materials,

It is possible to make mass production and reduce weight



f. Severe Plastic Deformation : Intense deformation at low temperatures (SPD methods)



FIGURE 2.4

equal channel angular extrusion (ECAE)

Schematic diagrams showing the (a) equal channel angular pressing (ECAP) and (b) highpressure torsion (HPT) processes.

HPT process: Equivalent strain induced on sample



HPT: severe plastic deformation, but Inhomogeneous process

f. Severe Plastic Deformation : Intense deformation at low temperatures (SPD methods)

• The SPD methods have some advantages over other methods in <u>synthesiz-</u> ing ultrafine-grained materials. Firstly, ultrafine-grained materials with highangle grain boundaries could be synthesized. Secondly, the samples are dense and there is no porosity in them since they have not been produced by the consolidation of powders. Thirdly, the grain size is fairly uniform throughout the structure. Lastly, the technique may be applied directly to commercial cast metals.

• However, the limitations of this technique are that the minimum grain size is not very small (typically it is about 200–300nm), and that amorphous phases or other metastable phases have not been synthesized frequently. But, these methods are very useful in producing bulk ultrafine-grained materials.

The methods of SPD described above have two drawbacks. Firstly, forming machines with large load capacities and expensive dies are required. Secondly, the productivity is relatively low.

g- Accumulative Roll Bonding (ARB process)

: 2 sheets of the same material are stacked, heated (to below the recrystallization temperature), and rolled, bonding the 2 sheets together. This sheet is cut in half, the 2 halves are stacked, and the process is repeated several times. Compared to other SPD processes, ARB has the benefit that it does not require specialized equipment. However, the surfaces to be joined must be well cleaned before rolling to ensure good bonding.



Schematic of the accumulative roll-bonding (ARB) process.