

September 27, 2014

Cheiron School 2014

X-ray Beamline Design 1  
**X-ray Monochromator**

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**SPring-8/JASRI**

# Outline

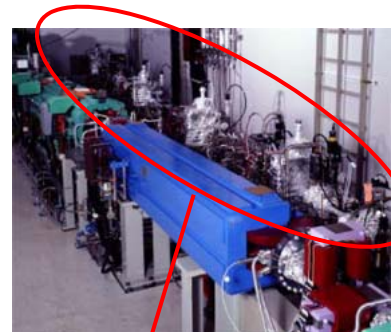
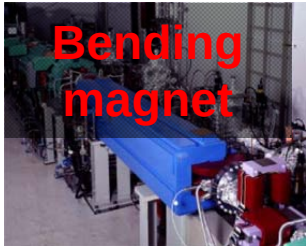
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1. Introduction
2. Light source
3. X-ray Monochromator
  - Fundamental of Bragg reflection
  - Dynamical theory
  - DuMond diagram ~ extraction of x-rays from SR
  - Double crystal monochromator
  - Crystal cooling
4. Summary

# Beamline structure

Beamline = “Bridge” between light source & experimental station

## Light source



## Front-end

## Experimental station

## Shielding hutch

## Ring tunnel

## Exp. hall



## Optics & transport

Monochromator, mirror  
shutter, slit  
pump,...

## → Transport and processing of photons

photon energy, energy resolution,  
beam size, beam divergence, polarization,...

## → Vacuum

protection of ring vacuum and beamline vacuum

## → Radiation safety

Shielding and interlock

# Light sources & X-ray optics

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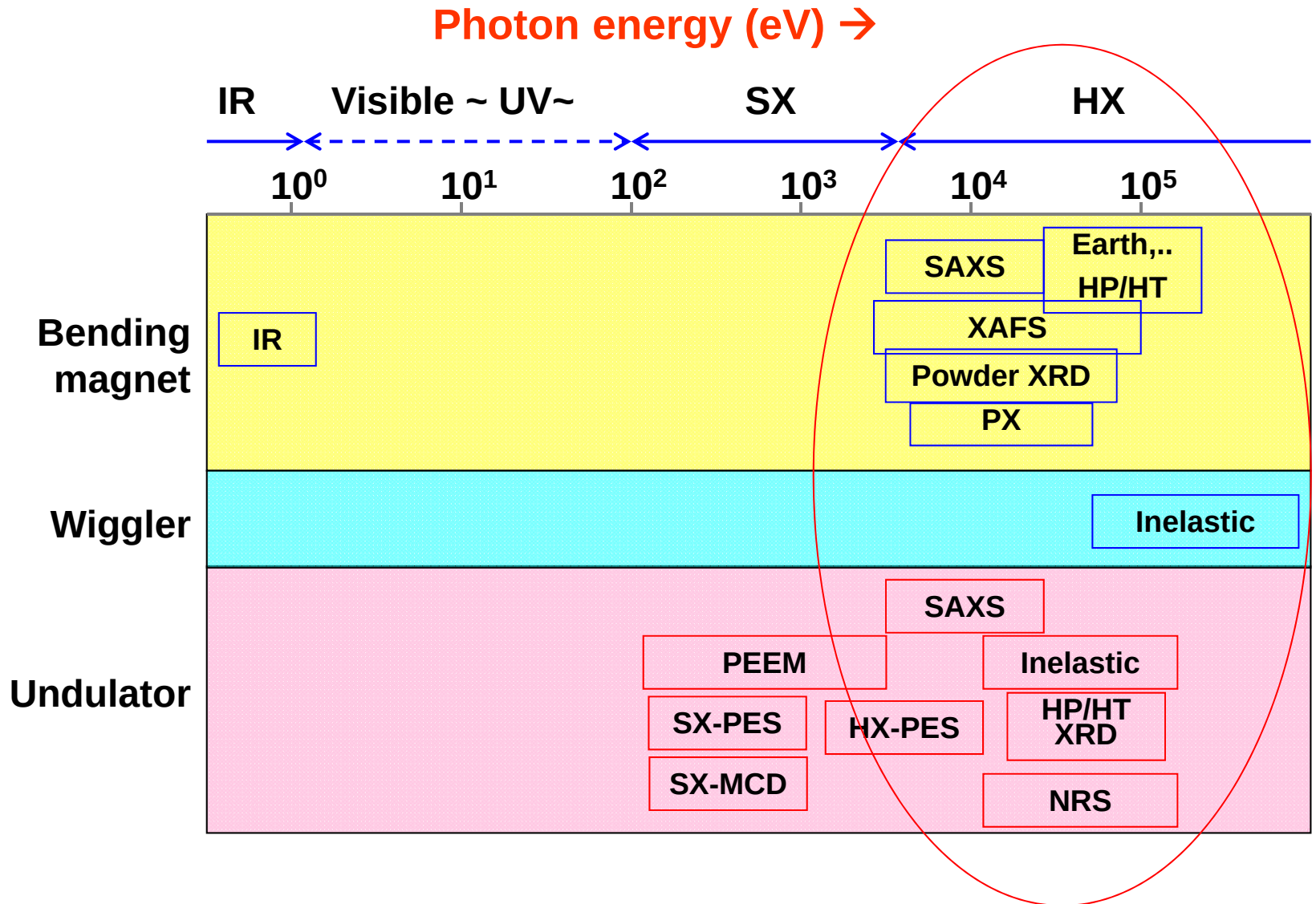
Check points to be considered for your SR application:

- White or monochromatic
- Energy range
- Energy resolution
- Flux & flux density
- Beam size at sample (micro beam?,...)
- Beam divergence/convergence at sample (Resolution in k-space)
- Higher order elimination w/ mirror
- Polarization conversion
- Spatial coherency

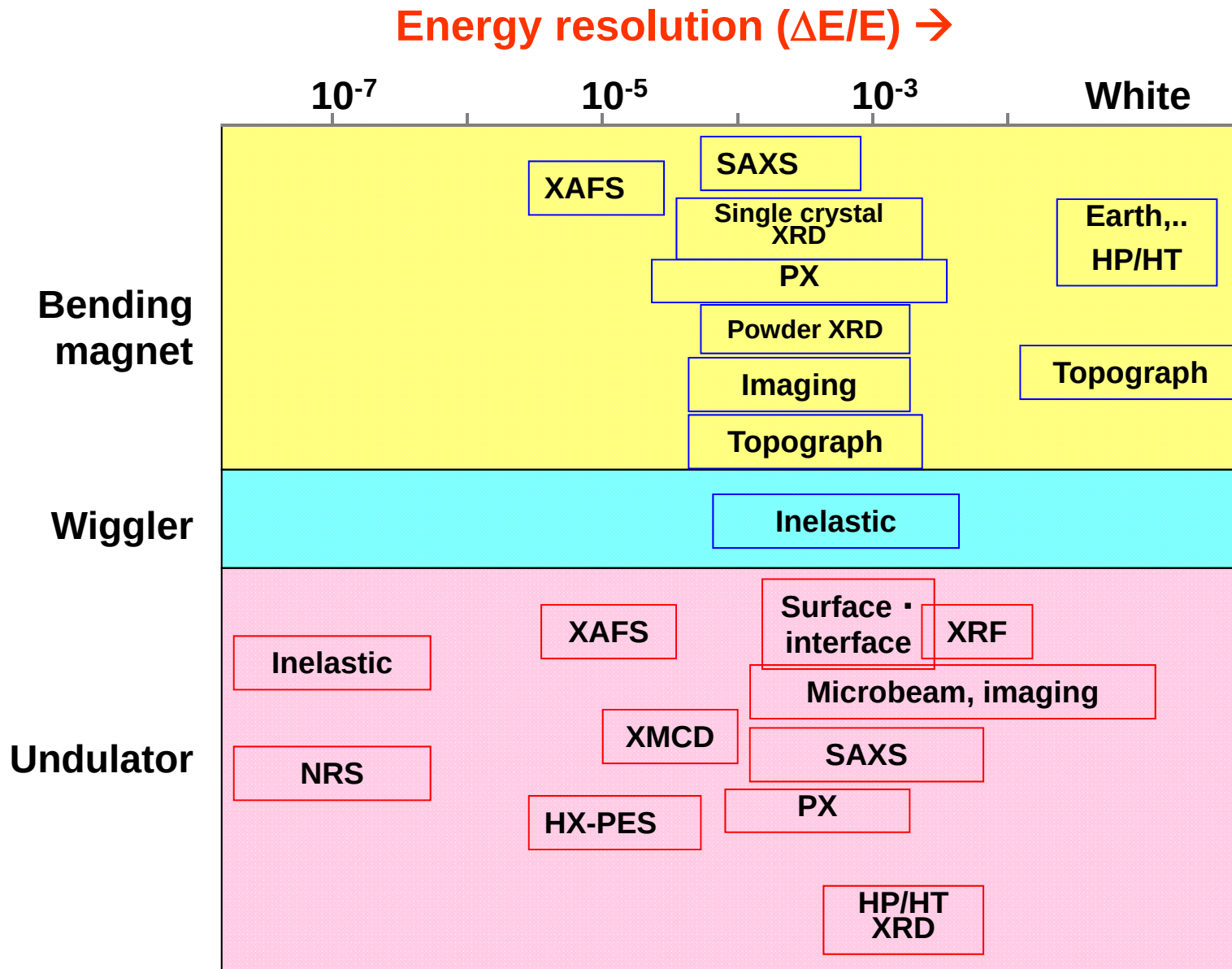
....

→ Light source, [monochromator](#), mirror,  
and other optical devices and components

# BL classification (energy region)



# BL classification (energy resolution)



# Light sources (1)

## Bending magnet or insertion devices ?

### Bending magnet:

- for wide energy range, continuous spectrum
- for wide beam application for large samples

### Undulator (major part of 3GLS beamline):

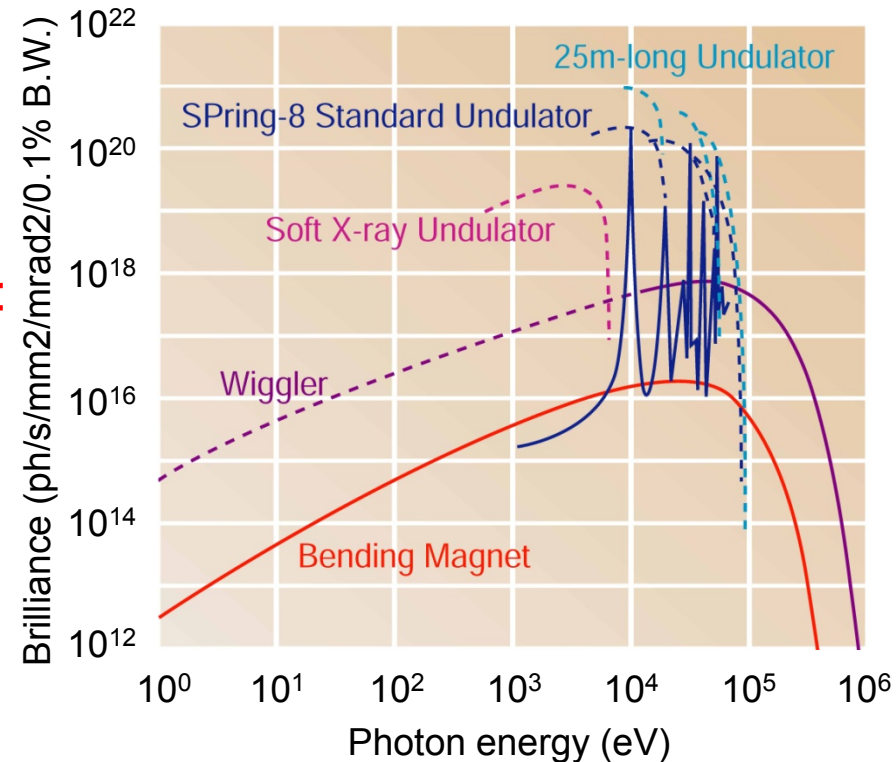
- for high-brilliance beam
- for micro-/ nano-focusing beam

### Wiggler:

- for higher energy X-rays > 100 keV.

Power, brilliance, flux density, partial flux,...  
can be calculated using code.

e.g. "SPECTRA" by T. Tanaka & H. Kitamura



Brilliance for SPRING-8 case

# Light sources (2)

Angular divergence and band width

→ Core part we need

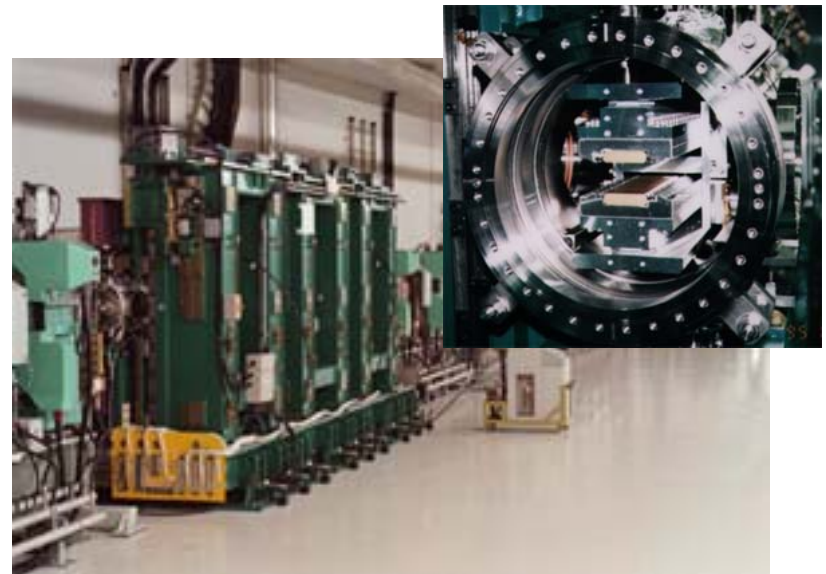
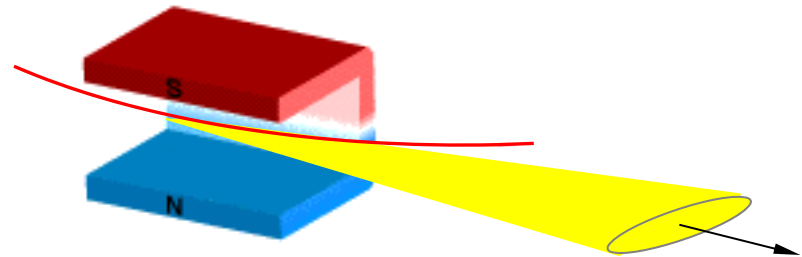
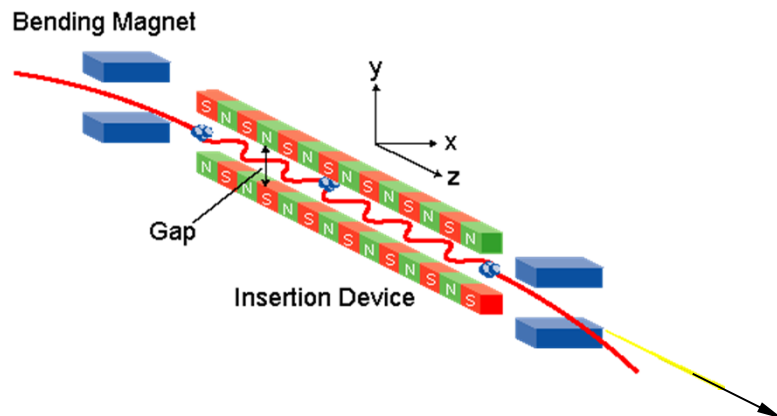
## Bending magnet

$$\sigma_{r'} \approx 0.597 \frac{1}{\gamma} \sqrt{\frac{\lambda}{\lambda_c}}$$

## Undulator

$$\sigma_{r'} \approx \sqrt{\frac{\lambda_n}{2N\lambda_u}} = \frac{1}{2\gamma} \sqrt{\frac{1+K^2/2}{nN}}$$

$$\frac{\Delta E}{E} \approx \frac{1}{nN}$$



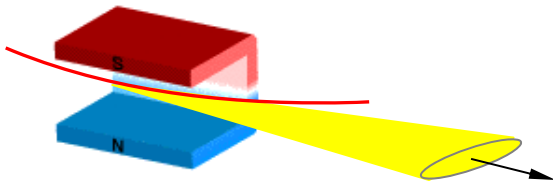
SPring-8 in-vacuum undulator



# Light sources (3)

Kilowatt of SR power → mostly eliminated before/by monochromator

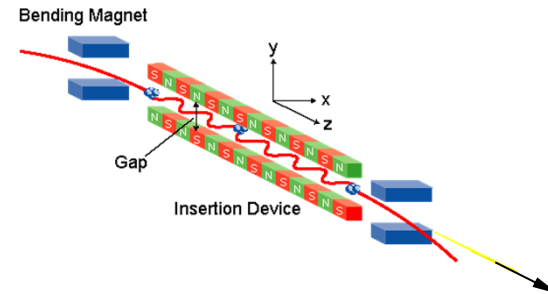
## Bending magnet



## Power distribution

$$\begin{cases} \psi_v \approx 1/\gamma \\ \psi_h \approx \text{const} \end{cases}$$

## Undulator



$$\begin{cases} \psi_v \approx 1/\gamma \\ \psi_h \approx K/\gamma \end{cases}$$

K: deflection parameter  
(K= 0.5~2.5)

## Total power

$$P_{\text{tot}} [\text{kW}] = 1.27 E^2 [\text{GeV}] B^2 [\text{T}] \overbrace{L_{\text{Arc}} [\text{m}]} R [\text{m}] \phi [\text{rad}] I [\text{A}]$$

$$E = 8 \text{ GeV}, I = 0.1 \text{ A}, B = 0.68 \text{ T}, R = 39.3 \text{ m} \\ \rightarrow P_{\text{tot}} = 0.15 \text{ kW/mrad}$$

$$P_{\text{tot}} [\text{kW}] = 1.27 E^2 [\text{GeV}] \frac{1}{2} B_0^2 [\text{T}] \overbrace{L [\text{m}]} \left\langle B_0^2 \sin^2(2\pi z / \lambda_u) \right\rangle I [\text{A}]$$

$$B_0 = 0.87 \text{ T}, L = 4.5 \text{ m} \\ \rightarrow P_{\text{tot}} = 14 \text{ kW}$$

# X-ray Monochromator

X-ray monochromator is key component for SR experiments:

- length gauge for structure analysis,
- energy gauge for spectroscopy,...

## Principle of x-ray monochromator

Photon energy tuning  $\leftarrow$  Bragg's law

Energy resolution  $\leftarrow$  source divergence, Darwin width,...

Flux (throughput)  $\leftarrow$  related to Darwin width

➔ Understanding the dynamical theory for large & perfect crystal

## Practical of the monochromator

- Double-crystal monochromator for fixed-exit

*Single-bounce monochromator is for limited case*

- Crystal cooling

High heat load depending on light source

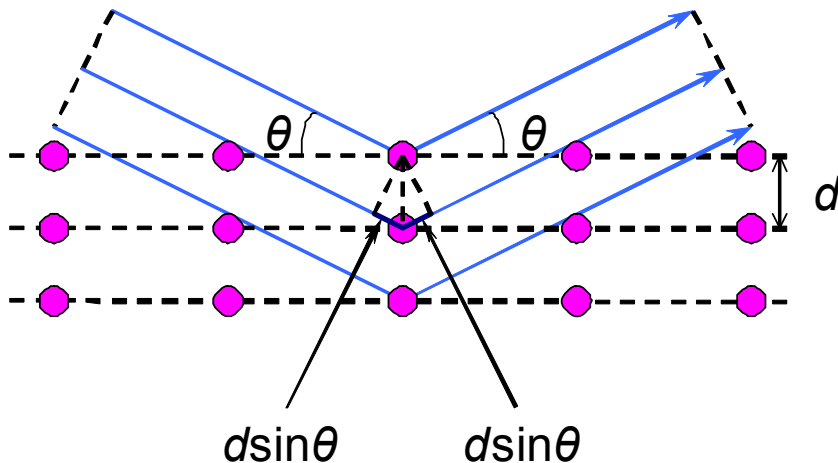
➔ Mechanical engineering issues

# Bragg reflection

## Bragg's law in real space

- 1) Phase matching on the single net plane by mirror-reflection condition.
- 2) Phase matching between net planes.

$$2d \sin \theta_B = m\lambda$$

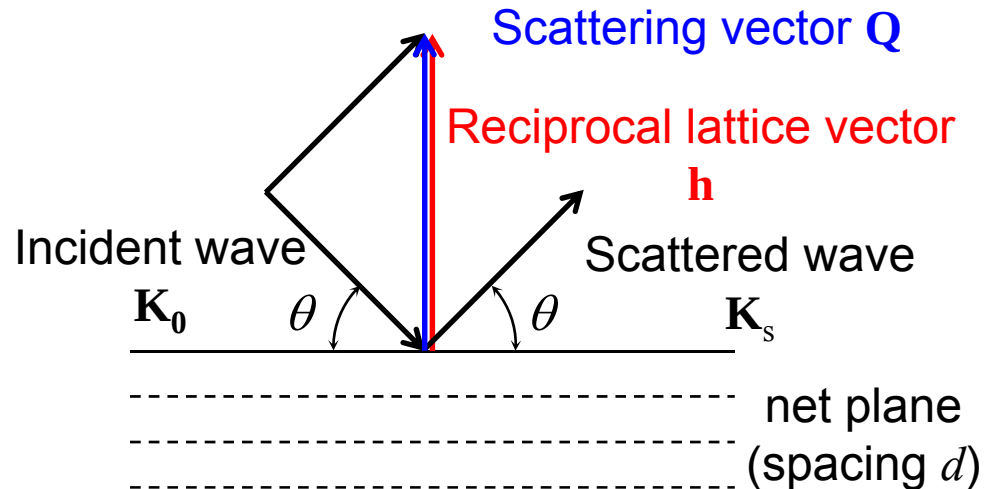


## Laue condition (Kinematical) in reciprocal space

$$\mathbf{Q} = \mathbf{K}_s - \mathbf{K}_0 = \mathbf{h}$$

Reciprocal lattice vector  $\mathbf{h}$

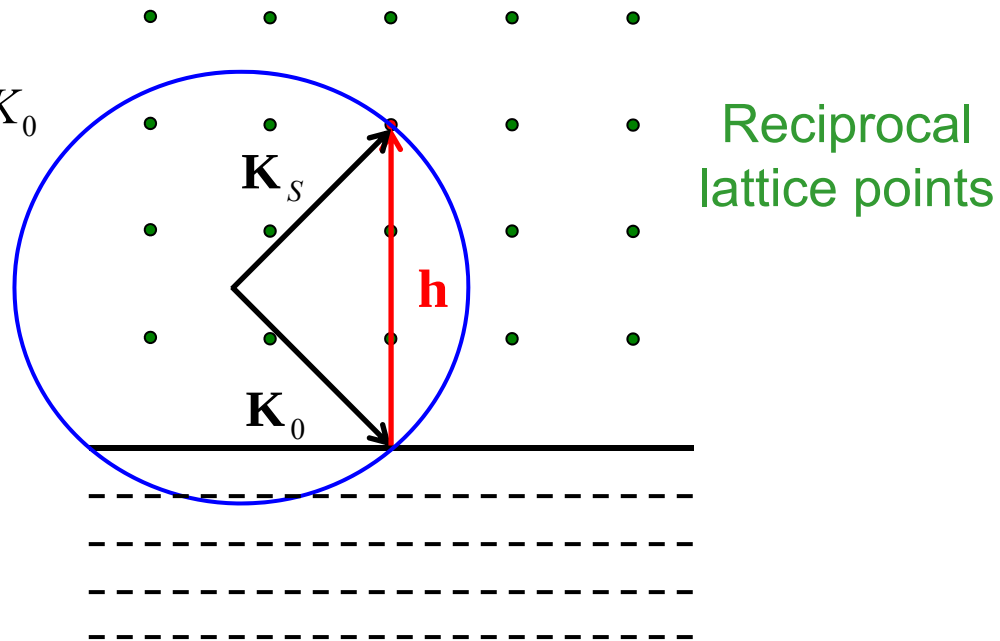
- Normal to net plane
- Length =  $1/d$



# Ewald sphere

Ewald sphere:

$$\text{Radius} = 1/\lambda = K_0$$

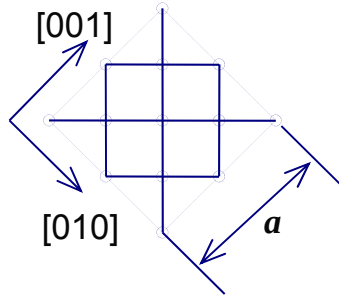


→ When a reciprocal lattice point is on the Ewald sphere, Bragg reflection occurs.

# Miller indices and $d$ -spacing for silicon

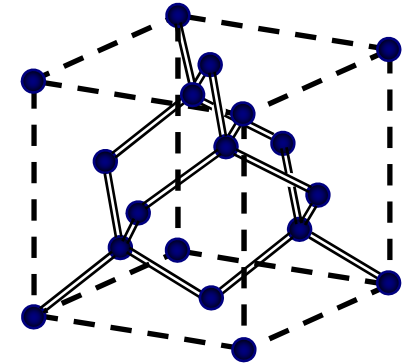
$$d = \frac{a}{\sqrt{h^2 + k^2 + l^2}}$$

Top view

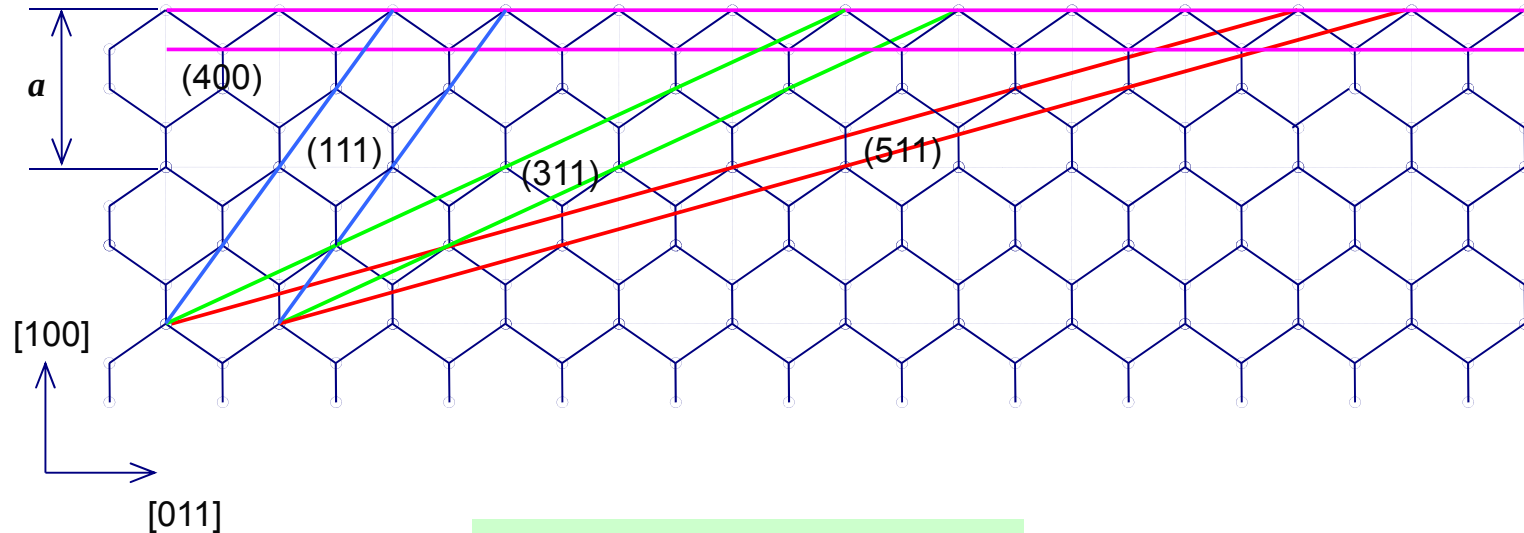


$$a = 5.431 \text{ \AA}$$

|              |                  |
|--------------|------------------|
| $d$ -spacing | (400) : 1.3578 Å |
|              | (111) : 3.1356 Å |
|              | (311) : 1.6375 Å |
|              | (511) : 1.0452 Å |



Side view



Diamond :  $a = 3.567 \text{ \AA}$

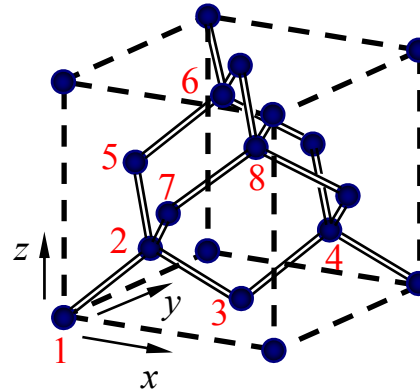
# Crystal structure factor for diamond structure

Structure factor  $\rightarrow$  Sum of atomic scattering with phase shift in the unit cell

$$F(\mathbf{h}) = \sum_j \underline{f_j(\mathbf{h}, E)} \exp(2\pi i \mathbf{h} \cdot \mathbf{r}_j)$$

Atomic scattering factor

$$F(\mathbf{h}) = \sum_j f_j(\mathbf{h}, E) \exp\{2\pi i(hx_j + ky_j + lz_j)\}$$



Position of atoms in the unit cell for diamond structure

$$(x_j, y_j, z_j) =$$

$$(0, 0, 0)_1, (1/4, 1/4, 1/4)_2,$$

$$(1/2, 1/2, 0)_3, (3/4, 3/4, 1/4)_4,$$

$$(0, 1/2, 1/2)_5, (1/4, 3/4, 3/4)_6,$$

$$(1/2, 0, 1/2)_7, (3/4, 1/4, 3/4)_8$$

For diamond structure

$$\begin{cases} h, k, l & \text{Mixture of odd and even numbers} \\ & F = 0 \\ h, k, l & \text{All odd, or, all even numbers, and } m: \text{integer,} \end{cases}$$

$$\begin{cases} h + k + l = 4m & F = 8f & \leftarrow 8 \text{ atoms in phase} \\ h + k + l = 4m \pm 1 & F = 4(1 \pm i)f & \leftarrow \text{Half contribute with phase shift } \pm \pi/2 \\ h + k + l = 4m \pm 2 & F = 0 & \leftarrow \text{Half cancel with } \pi \end{cases}$$

# Crystal structure factor for diamond structure

(400), (220),...

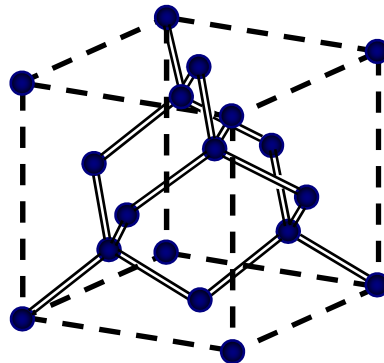
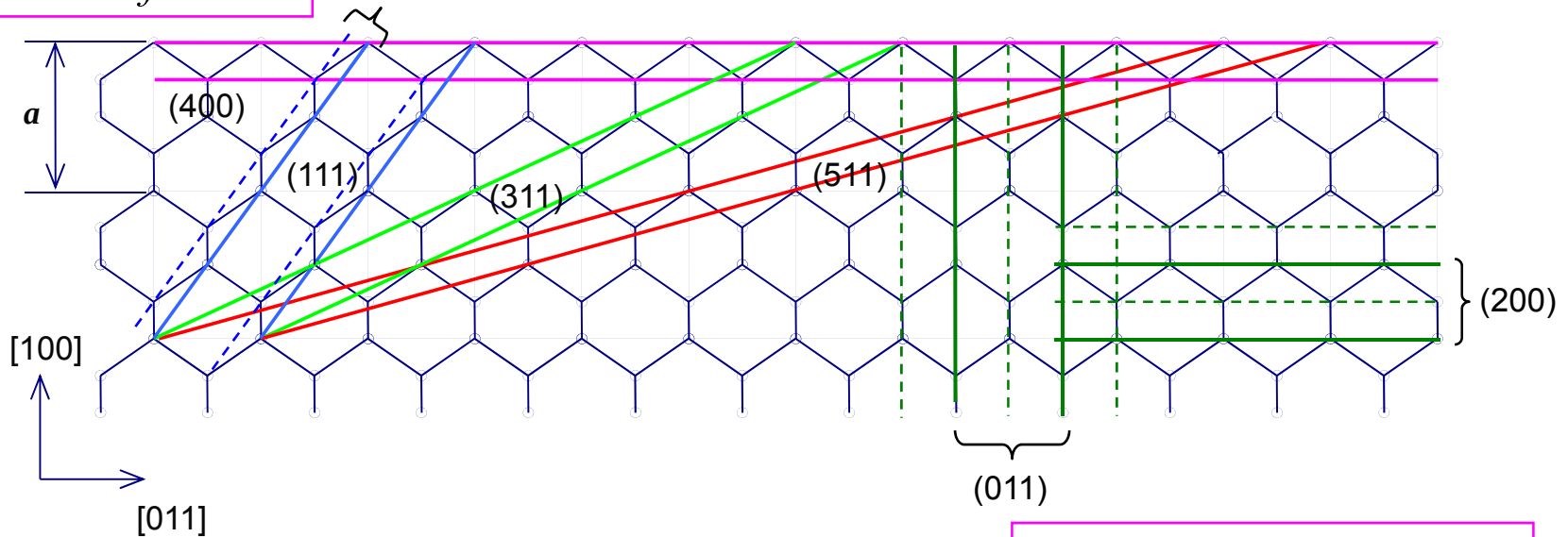
All in phase

$$\rightarrow F = 8f$$

(111), (311),...

Half contribute with phase shift  $\pm \pi/2$

$$\rightarrow F = 4(1 \pm i)f$$



(011), (200),...

Half cancel with  $\pi$

$\rightarrow$  Forbidden reflection

$$F = 0$$

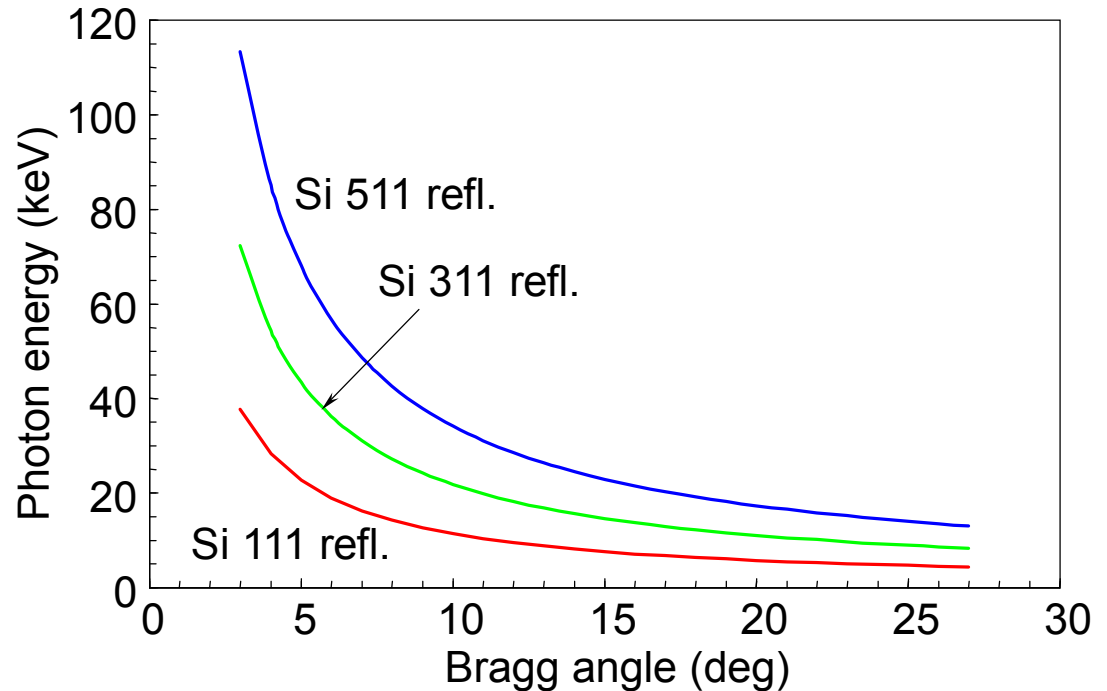
# X-ray monochromator using perfect crystal

→ Perfect single crystal: **silicon, diamond**,..

Photon energy tuning:

- Crystal & lattice plane
- Bragg angle range

$$E [\text{keV}] = \frac{12.3984}{2d_{hkl} [\text{\AA}] \sin \theta_B}$$



e.g. for SPring-8 standard DCM

Bragg angle: 3~27°



# Kinematical X-ray diffraction

3-dimensional periodic structure of unit cell with number  $N_x, N_y, N_z$

Total scattering intensity becomes:

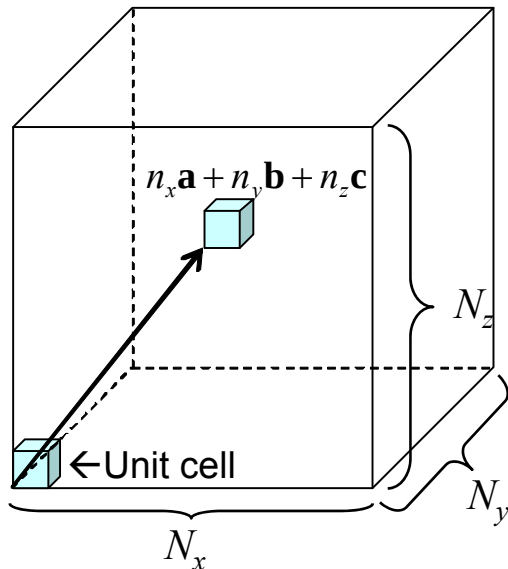
$$I = I_e |F(\mathbf{Q})|^2 \cdot |G(\mathbf{Q})|^2$$

Laue function:  $|G(\mathbf{Q})|^2 = \frac{\sin^2(\pi N_x h)}{\sin^2(\pi h)} \cdot \frac{\sin^2(\pi N_y k)}{\sin^2(\pi k)} \cdot \frac{\sin^2(\pi N_z l)}{\sin^2(\pi l)}$

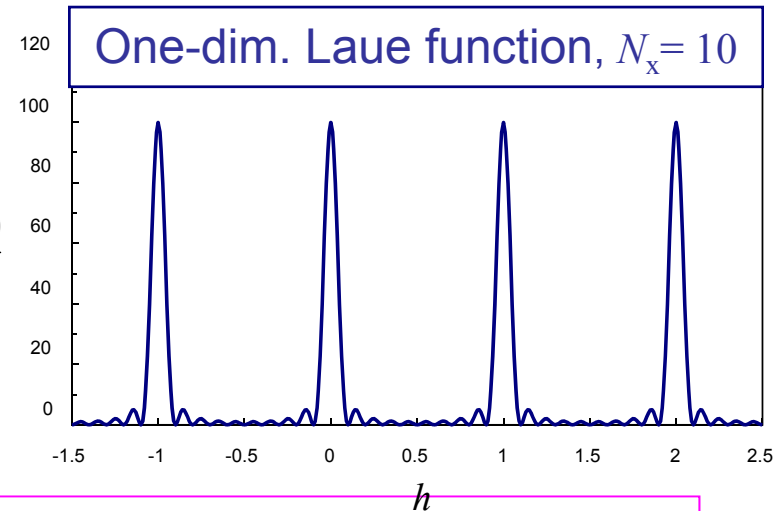
← 3-dim. Periodic structure

$h, k, l$ : integer → Intense peaks

→  $(hkl)$  reflection



$$\frac{\sin^2(\pi N_x h)}{\sin^2(\pi h)}$$



Peak intensity  $N_x^2$

FWHM  $\Delta h \approx 0.8858/N_x \sim 1/N_x$

Crystal size becomes larger → narrower & higher, approaching delta function

# **Dynamical theory**

## ***Two-beam approximation***

# Kinematical to dynamical theory

“Large & perfect” single crystal:

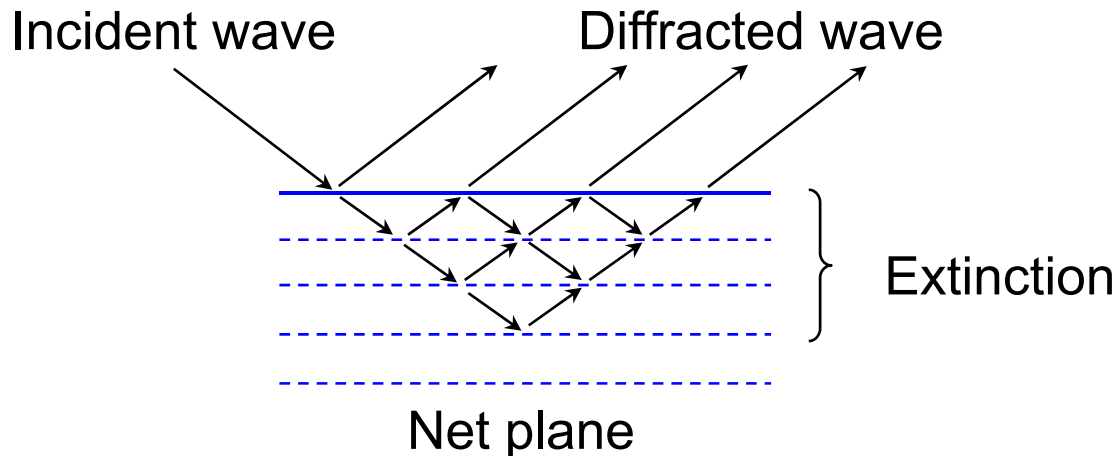
1) Multiple scattering w/  $\mathbf{h}$  &  $-\mathbf{h}$  reflection

2) Extinction

(Diffraction by “finite” number of net planes)

Kinematical diffraction is invalid

→ Dynamical theory must be applied.



# Fundamental equation

Fundamental equation is derived

using **Maxwell's equations** and introducing **Bloch wave**

for 3-dimensional periodic medium (= perfect single crystal):

$$\frac{k_h^2 - K_0^2}{K_0^2} E_h = \sum_g \chi_{h-g} (\mathbf{e}_h \cdot \mathbf{e}_g) E_g$$

$h, g, \dots$  : Reciprocal lattice points

$E_h, E_g$  : Fourier components of electric field

$K_0$  : Incident wave vector in vacuum

$k_h$  : Wave vectors in the crystal

$\chi_h$  : Fourier components of the polarizability (Negative values,  $10^{-6} \sim 10^{-5}$ )

$P = (\mathbf{e}_h \cdot \mathbf{e}_g)$  : Polarization factor between  **$h$**  and  **$g$**  waves

$k_h = k_0 + h$  : Momentum conservation

# Two-beam approximation

Fundamental equation is reduced to the equation for  
**two beams (waves)** of incidence and “one” intense diffraction

$$\frac{k_h^2 - K_0^2}{K_0^2} E_h = \sum_g \chi_{h-g} (\mathbf{e}_h \cdot \mathbf{e}_g) E_g$$



$$(A) \quad \frac{k_0^2 - K_0^2}{K_0^2} E_0 = \chi_0 E_0 + P \chi_{-h} E_h$$

$$(B) \quad \frac{k_h^2 - K^2}{K^2} E_h = P \chi_h E_0 + \chi_0 E_h$$

$\chi_0, \chi_h, \chi_{-h}$  : Fourier components of the polarizability

(Negative values,  $10^{-6} \sim 10^{-5}$ )

$P = (\mathbf{e}_0 \cdot \mathbf{e}_h)$  : Polarization factor ( $\sigma : P = 1, \pi : P = \cos 2\theta_B$ )

# Two-beam approximation

Using two equations, we obtain following secular equation:

$$(A) \quad \frac{k_0^2 - K_0^2}{K_0^2} E_0 = \chi_0 E_0 + P \chi_{-h} E_h$$

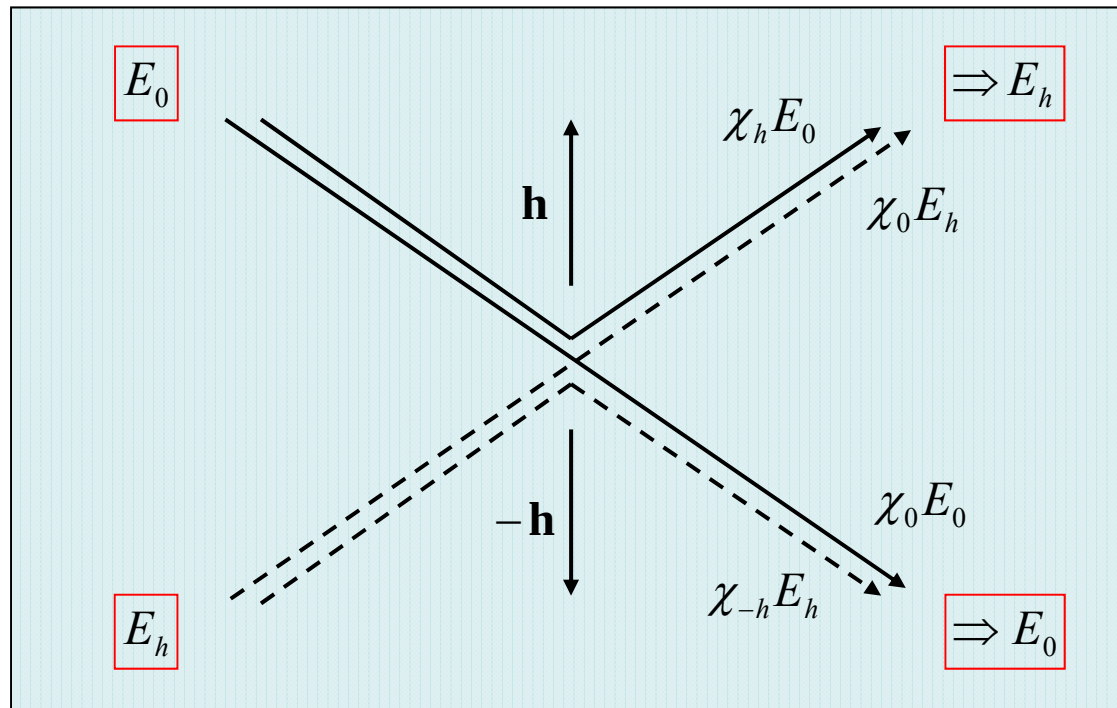
$$(B) \quad \frac{k_h^2 - K^2}{K^2} E_h = P \chi_h E_0 + \chi_0 E_h$$

Secular equation

$$(\mathbf{k}_0^2 - k^2)(\mathbf{k}_h^2 - k^2) = \chi_h \chi_{-h} P^2 K_0^4$$

$$k^2 = (1 + \chi_0) K_0^2$$

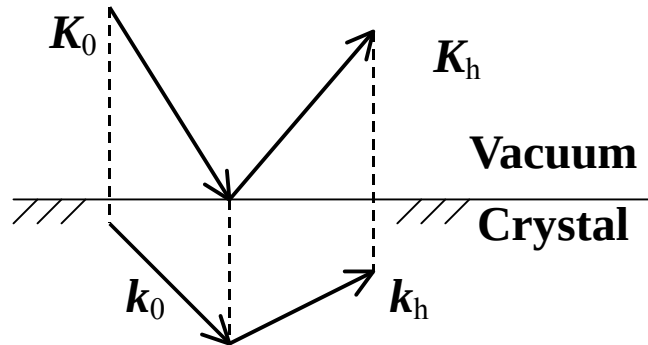
$k$  : Mean wave number in the crystal



Scheme of self-consistent wave field

# Boundary condition of wave vector

We must consider connections of waves from vacuum into the crystal and from the crystal to vacuum, to solve the equations.



**Tangential component of wave vector must be continuous.**

Incident wave in vacuum

→ Refracted wave in the crystal

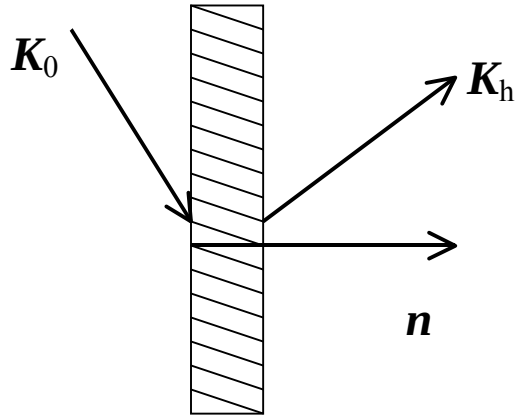
→ Bragg reflection in the crystal

→ Reflected wave in the crystal

→ Reflected wave in vacuum

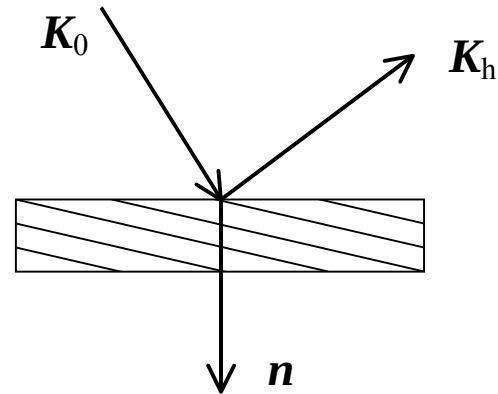
# Laue case and Bragg case

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**Laue case**

(Transmission geometry)



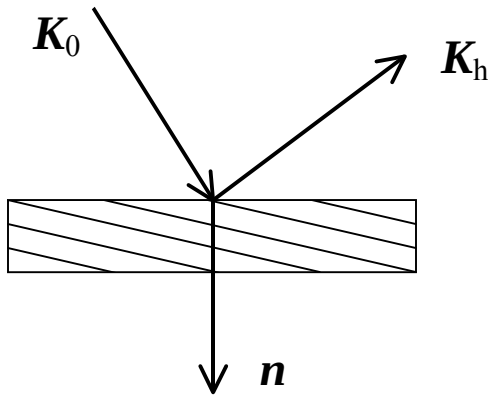
**Bragg case**

(Reflection geometry)



# Asymmetry ratio

$$\begin{cases} \gamma_0 = \hat{K}_0 \cdot n \\ \gamma_h = \hat{K}_h \cdot n \end{cases} \quad b = \frac{\gamma_0}{\gamma_h}$$



**Laue case:  $b > 0$**

Symmetric Laue case:  $b = 1$

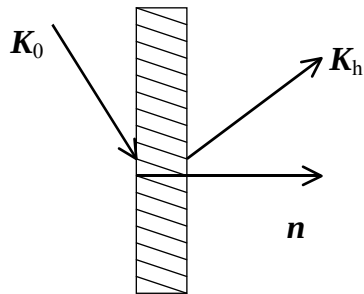
**Bragg case:  $b < 0$**

Symmetric Bragg case:  $b = -1$

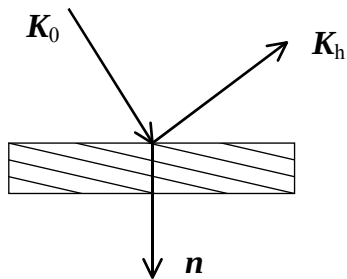
$n$ : normal vector to the surface

# Dispersion surface

$(\mathbf{k}_0^2 - k^2)(\mathbf{k}_h^2 - k^2) = \chi_h \chi_{-h} P^2 K_0^4$  ← Secular equation is **quartic** equation, and it gives four-point solution on the  $\mathbf{n}$ -vector, producing the **dispersion surfaces**.

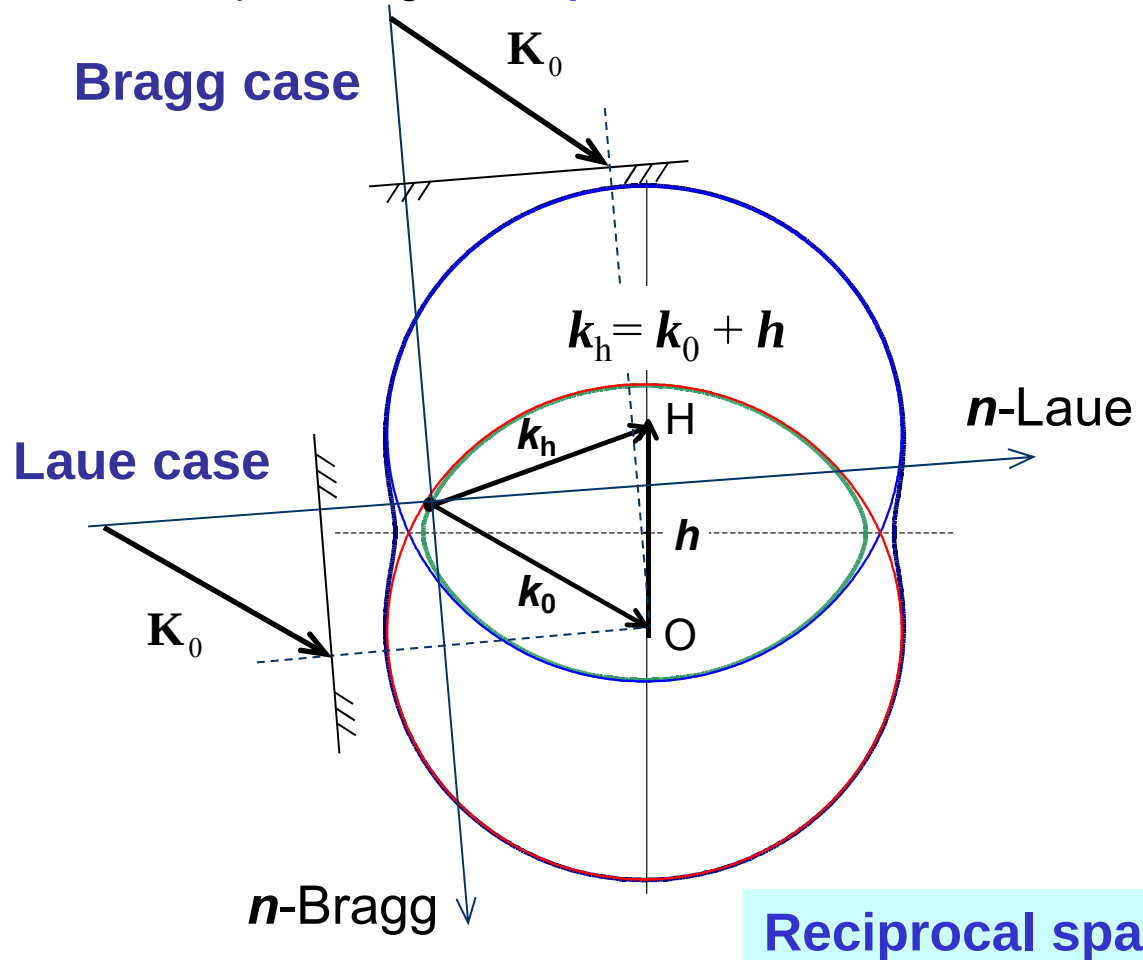


Laue case



Bragg case

Real space



Reciprocal space

Two dispersion surfaces show the gap near Bragg condition.

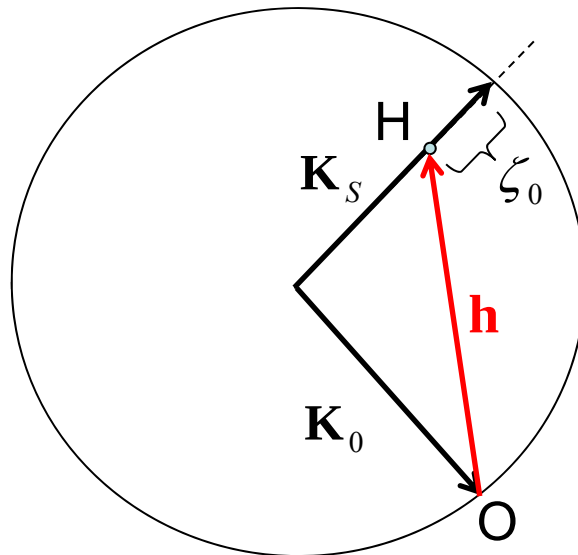
# Deviation from Bragg condition

## Excitation error

→ Geometrical deviation  $\zeta_0$  from Bragg condition:

Distance between Ewald sphere and the reciprocal lattice point.

$\zeta_0$  is positive when H is inside the Ewald sphere (by S. Miyake).

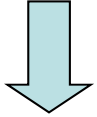


$$\zeta_0 \approx -\frac{2(\mathbf{K}_0 \cdot \mathbf{h}) + h^2}{2K_0}$$

# Normalized deviation parameter $W$

Parameter  $W$  is related to the gap between two dispersion surfaces and total reflection occurs at  $-1 < W < 1$  for Bragg case.

$$W = - \frac{2(\mathbf{K}_0 \cdot \mathbf{h}) + h^2}{2K_0^2} \sqrt{\frac{\gamma_0}{|\gamma_h|}} \frac{1}{|\chi_{hr}| \cdot |P|} + \frac{\chi_{0r}}{2|\chi_{hr}| \cdot |P|} \sqrt{\frac{\gamma_0}{|\gamma_h|}} \left(1 - \frac{\gamma_h}{\gamma_0}\right)$$



$\Delta\theta$ : Angle deviation for fixed photon energy,

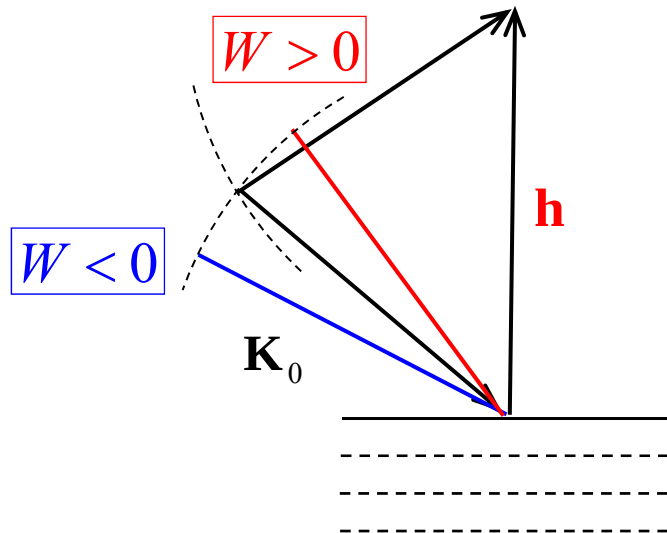
$\Delta E$ : Energy deviation for fixed incident angle

$$W = \left\{ \Delta\theta \sin 2\bar{\theta}_{BK} + 2 \frac{\Delta E}{E} \sin^2 \bar{\theta}_{BK} + \frac{\chi_{0r}}{2} \left(1 - \frac{\gamma_h}{\gamma_0}\right) \right\} \sqrt{\frac{\gamma_0}{|\gamma_h|}} \frac{1}{|\chi_{hr}| \cdot |P|}$$

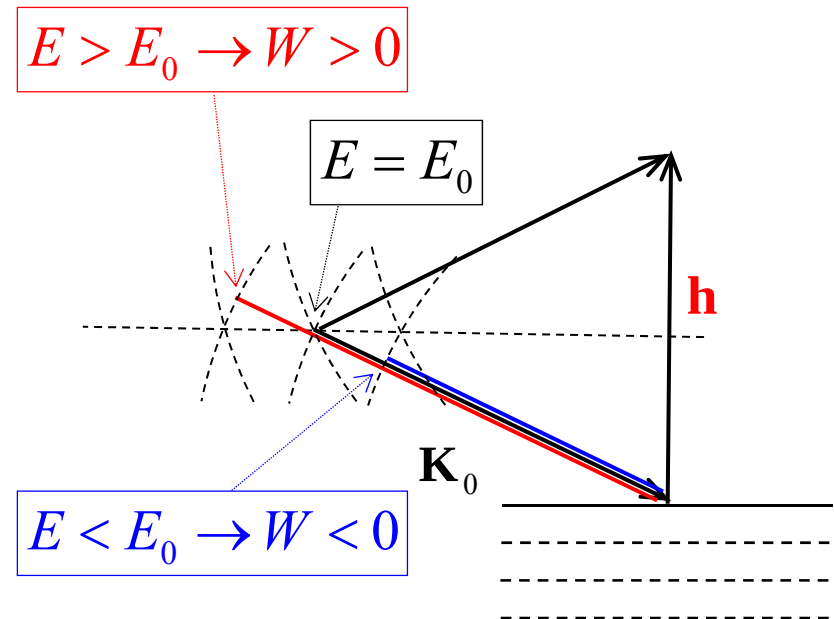
For symmetric Bragg case, sigma polarization:

$$W = \left\{ \Delta\theta \sin 2\bar{\theta}_{BK} + 2 \frac{\Delta E}{E} \sin^2 \bar{\theta}_{BK} + \chi_{0r} \right\} \frac{1}{|\chi_{hr}|}$$

# Sign of deviation parameter $W$



Angle deviation at fixed energy  
→ direction change of wave vector



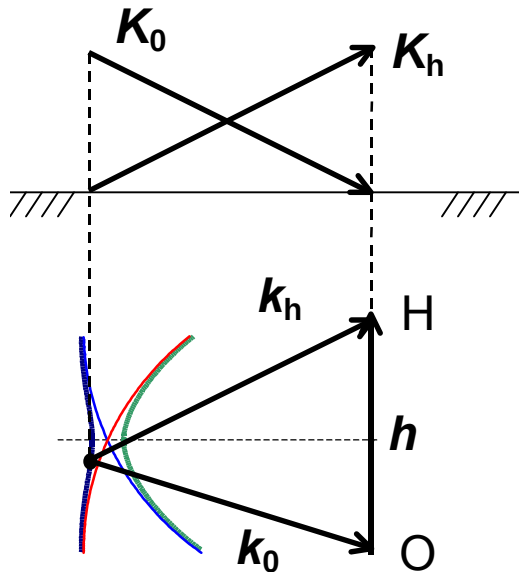
Energy deviation at fixed angle  
→ length change of wave vector

# Movement of tie point

Tie point moves by changing the incident angle  
at fixed photon energy (wavelength).

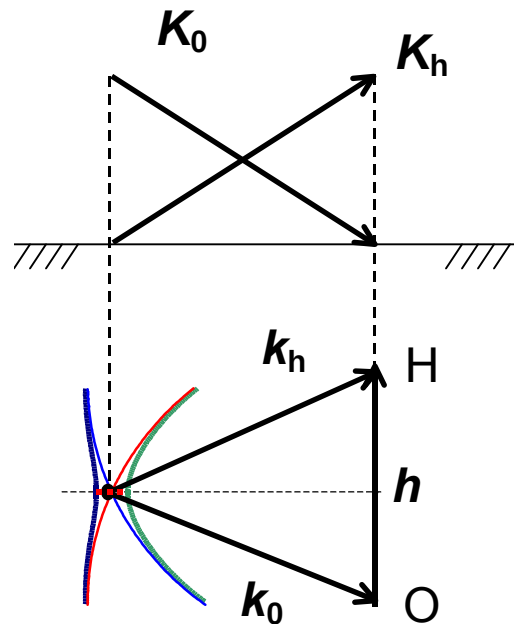
(1) Lower angle

$$W < -1$$



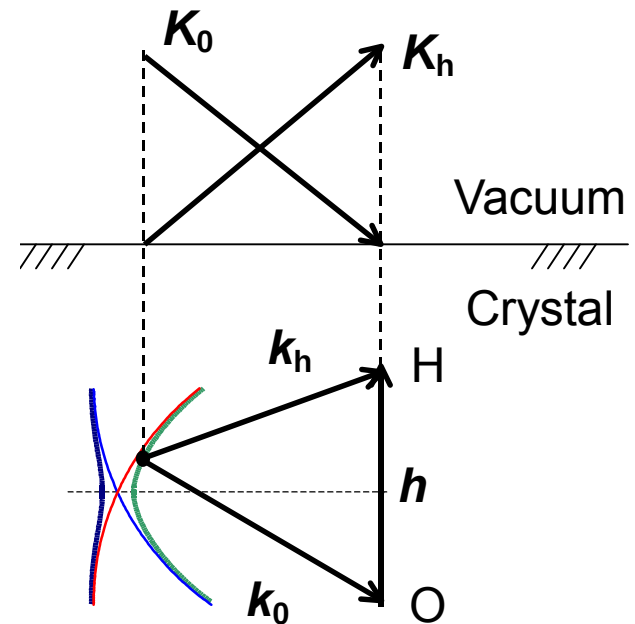
(2) Near Bragg condition

$$-1 < W < 1$$



(3) Higher angle

$$W > 1$$



**Total reflection**

Dominant branch for thick Bragg-case crystal is close to **O-sphere**.

# Calculation of polarizability

$\chi_h$ : Fourier component of polarizability  
 $\rightarrow$  proportional to the structure factor

$$\chi_h = -\frac{r_e \lambda^2}{\pi v_c} F(\mathbf{h}, E)$$

$v_c$ : unit cell volume

$$\chi_h = \chi_{hr} + i\chi_{hi}$$

$$\chi_{hr} \Leftrightarrow f^0(\mathbf{h}) + f'(E)$$

Atomic form factor  
 + real part of anomalous factor

$$\chi_{hi} \Leftrightarrow f''(E)$$

Imaginary part of  
 anomalous factor

For diamond structure

$$h + k + l = 4m$$

$$\chi_{hr} = -\frac{r_e \lambda^2}{\pi v_c} 8(f^0 + f')e^{-M}$$

$$\chi_{hi} = -\frac{r_e \lambda^2}{\pi v_c} 8f''e^{-M}$$

$$h + k + l = 4m \pm 1$$

$$\chi_{hr} = -\frac{r_e \lambda^2}{\pi v_c} 4(1+i)(f^0 + f')e^{-M}$$

$$\chi_{hi} = -\frac{r_e \lambda^2}{\pi v_c} 4(1+i)f''e^{-M}$$

$$h = k = l = 0$$

$$\chi_{0r} = -\frac{r_e \lambda^2}{\pi v_c} 8(Z + f')$$

$$\chi_{0i} = -\frac{r_e \lambda^2}{\pi v_c} 8f''$$

# Amplitude ratio

From the solution of the fundamental equations,  
we obtain the ratio  $r = E_h/E_0$  ( $\leftarrow$  reflection coefficient)  
as a function of parameter  $W$ .

For Bragg case, no absorption, and thick crystal:

$$\left\{ \begin{array}{ll} r = \frac{E_h}{E_0} = -\sqrt{\frac{\gamma_0}{|\gamma_h|}} \frac{|\chi_{hr}|}{\chi_{-h}} \frac{|P|}{P} \left( W + \sqrt{W^2 - 1} \right) & (W < -1) \\ \\ r = \frac{E_h}{E_0} = -\sqrt{\frac{\gamma_0}{|\gamma_h|}} \frac{|\chi_{hr}|}{\chi_{-h}} \frac{|P|}{P} \left( W + i\sqrt{1 - W^2} \right) & (-1 \leq W \leq 1) \quad \leftarrow \text{Total reflection} \\ \\ r = \frac{E_h}{E_0} = -\sqrt{\frac{\gamma_0}{|\gamma_h|}} \frac{|\chi_{hr}|}{\chi_{-h}} \frac{|P|}{P} \left( W - \sqrt{W^2 - 1} \right) & (W > 1) \end{array} \right.$$



# Reflectivity (Darwin curve)

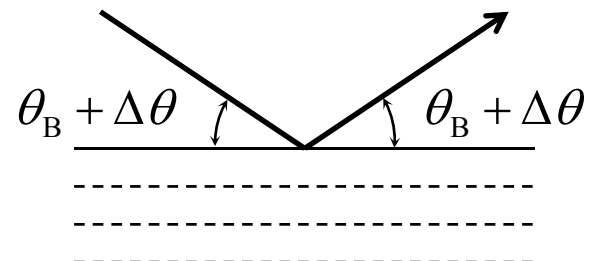
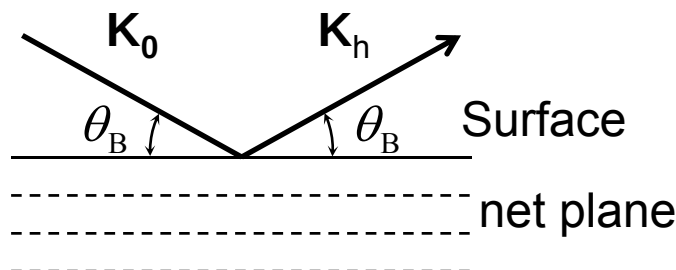
**Darwin curve** (intrinsic reflection curve for monochromatic plane wave)  
for Bragg case, no absorption, and thick crystal:

$$\begin{cases} R = \left(W + \sqrt{W^2 - 1}\right)^2 & (W < -1) \\ R = 1 & (-1 \leq W \leq 1) \quad \leftarrow \text{Total reflection region} \\ R = \left(W - \sqrt{W^2 - 1}\right)^2 & (W > 1) \end{cases}$$

$W$ : deviation parameter for s-polarization, symmetrical Bragg case

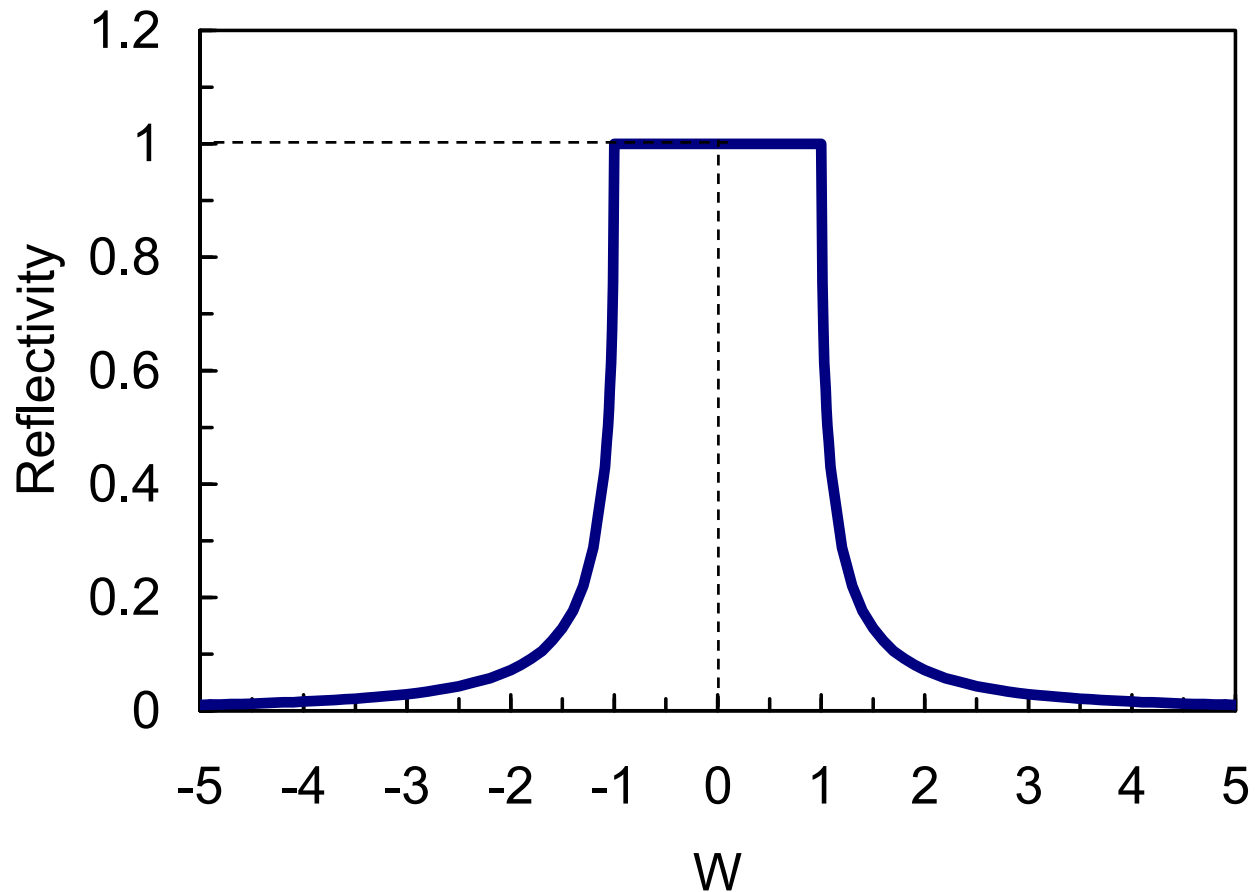
$$W = \left( \underbrace{\Delta\theta \sin 2\theta_B}_{\text{Angular deviation}} + \underbrace{2 \sin^2 \theta_B \frac{\Delta E}{E}}_{\text{Energy deviation}} + \underbrace{\chi_0}_{\text{Refraction}} \right) \frac{1}{|\chi_h|}$$

Geometry for  
symmetrical  
Bragg case



# Darwin curve

For Bragg case, **no absorption**, and thick crystal:



# Reflectivity with absorption

## Reflectivity

- symmetrical Bragg case,
- s-polarization,
- thick crystal

$$R = L - \sqrt{L^2 - 1}$$

$$L = \frac{\left\{ W^2 + g^2 + \sqrt{(W^2 - g^2 - 1 + \kappa^2)^2 + 4(gW - \kappa)^2} \right\}}{1 + \kappa^2}$$

$$W = \left( \Delta\theta \sin 2\bar{\theta}_B + 2 \sin^2 \bar{\theta}_B \frac{\Delta E}{E} + \chi_{0r} \right) \frac{1}{|\chi_{hr}|}$$

$$g = \frac{\chi_{0i}}{|\chi_{hr}|}, \quad \kappa = \frac{|\chi_{hi}|}{|\chi_{hr}|}$$

Note: No absorption  $g = 0, \kappa = 0 \Rightarrow R \rightarrow \text{Darwin curve}$

# Reflectivity curve for silicon

Examples for symmetrical Bragg case, **with absorption**,  
s-polarization and thick crystal:

## Si 111 refl., 10 keV

$$\chi_{0r} = -9.78 \times 10^{-6}$$

$$\chi_{0i} = -1.48 \times 10^{-7}$$

$$\chi_{111_r} = -3.66 \times 10^{-6}(1+i)$$

$$\chi_{111_i} = -7.30 \times 10^{-8}(1+i)$$

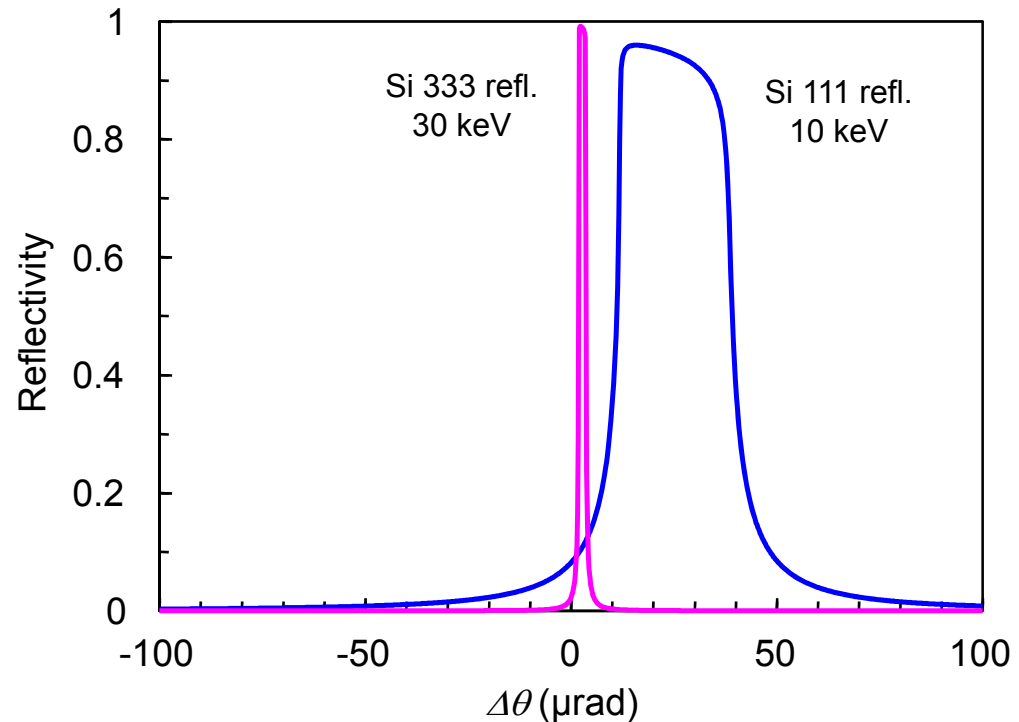
## Si 333 refl., 30 keV

$$\chi_{0r} = -1.07 \times 10^{-6}$$

$$\chi_{0i} = -1.75 \times 10^{-9}$$

$$\chi_{333_r} = -2.24 \times 10^{-7}(1+i)$$

$$\chi_{333_i} = -7.87 \times 10^{-10}(1+i)$$



- Width of  $0.1 \sim 100 \mu\text{rad}$
- Peak  $\sim 1$  with small absorption

# DuMond (angle-energy) diagram

The diagram helps to understand how we can extract x-rays from SR source.

Angular width  
(Darwin width)

$$\Delta\theta_{\text{Darwin}} = \frac{2|\chi_{hr}|}{\sin 2\theta_B} \propto |F(\mathbf{h})| \quad \leftarrow \Delta W = 2$$

Energy resolution

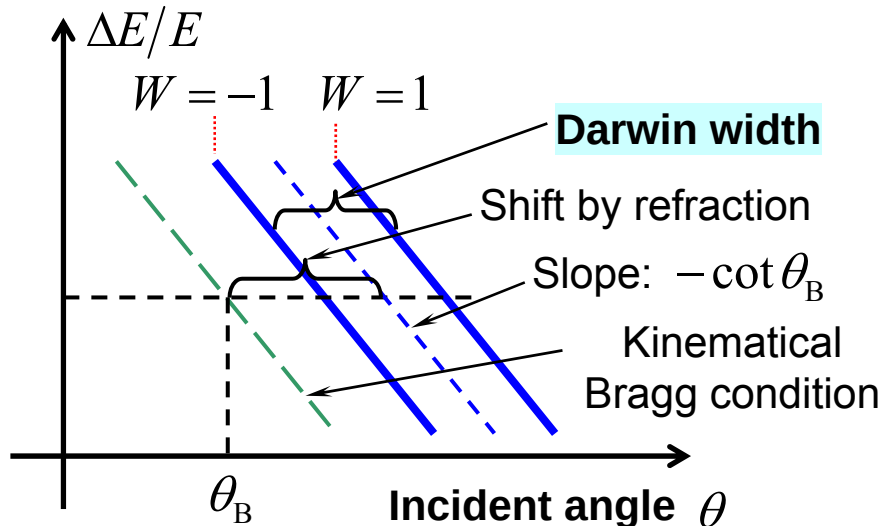
$$\frac{\Delta E}{E} = \cot \theta_B \sqrt{\Omega^2 + \Delta\theta_{\text{Darwin}}^2}$$

← Gaussian approximation for both light source and reflection curve

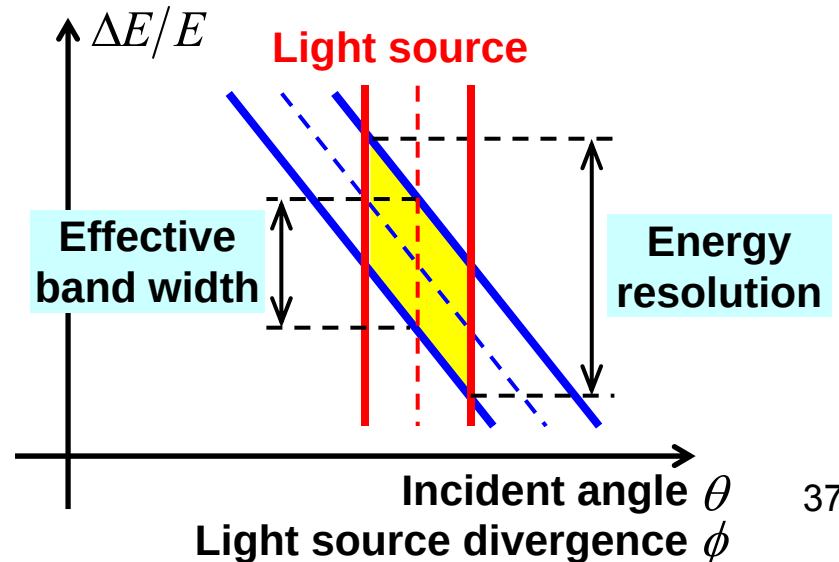
Effective band width

$$\frac{\Delta E}{E} \approx \frac{|\chi_{hr}|}{\sin^2 \theta_B}$$

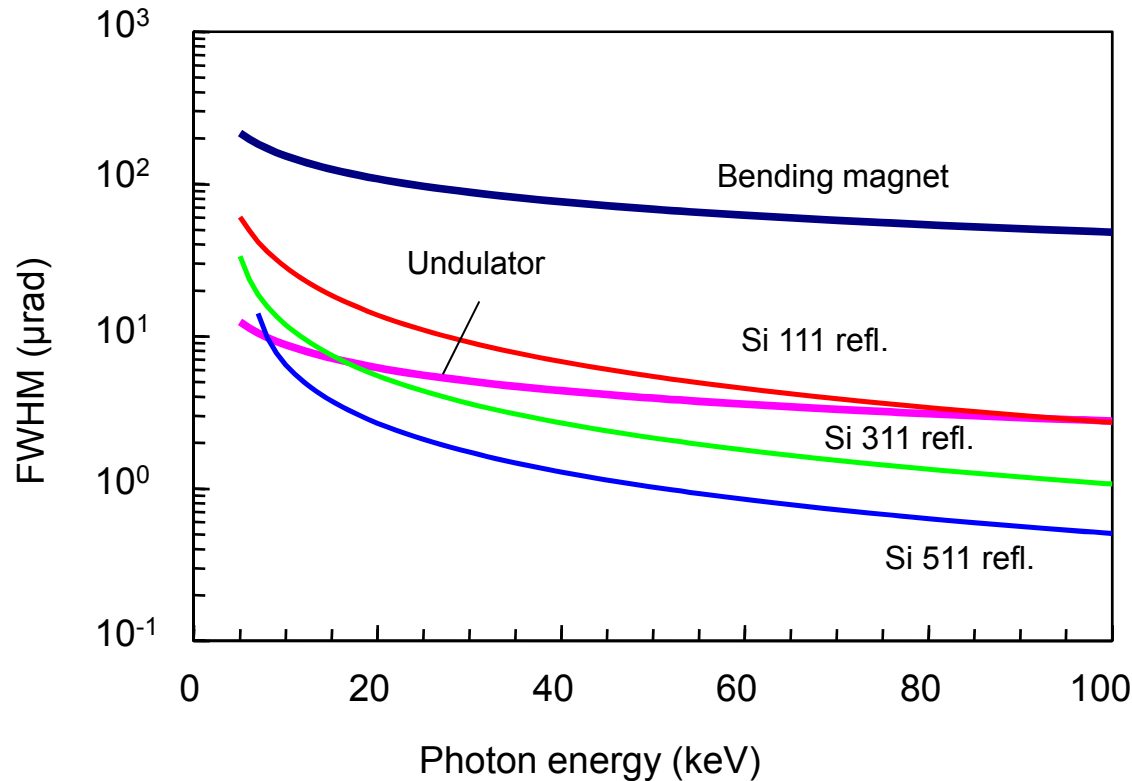
Relative energy



Relative energy



# Source divergence and diffraction width



Natural divergence

- Bending magnet

$$\sigma_{r'} \approx 0.597 \frac{1}{\gamma} \sqrt{\frac{\lambda}{\lambda_c}} \propto \sqrt{\frac{1}{\hbar\omega}}$$

- Undulator

$$\sigma_{r'} \approx \sqrt{\frac{\lambda}{2N\lambda_u}} \propto \sqrt{\frac{1}{\hbar\omega}}$$

For SPring-8 case:

- Bending magnet

$$\sigma_{r'} \approx 60 \mu\text{rad}$$

- Undulator ( $N=140$ )

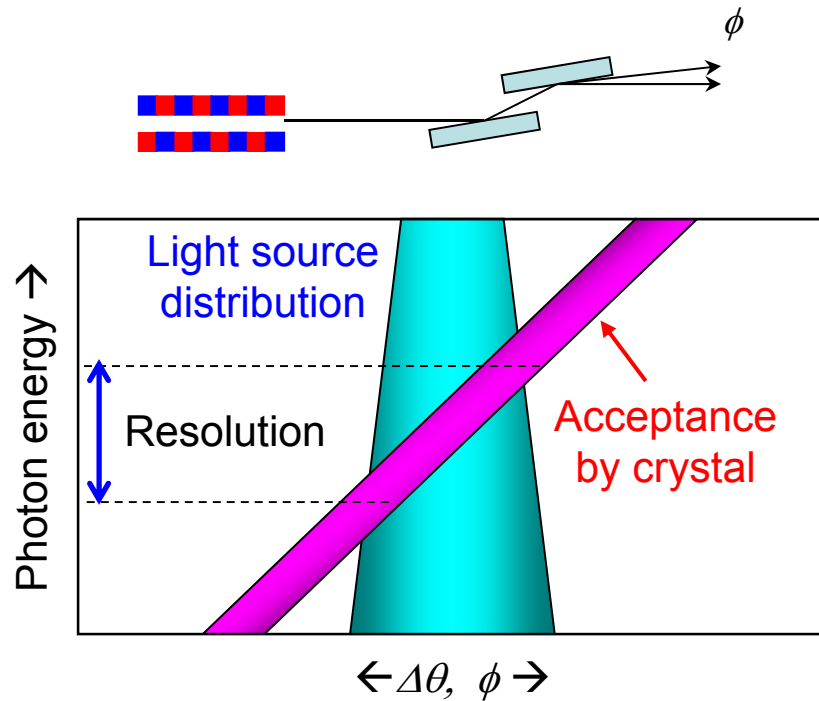
$$\sigma_{r'} \approx 5 \mu\text{rad}$$

Divergence of undulator radiation ~ diffraction width

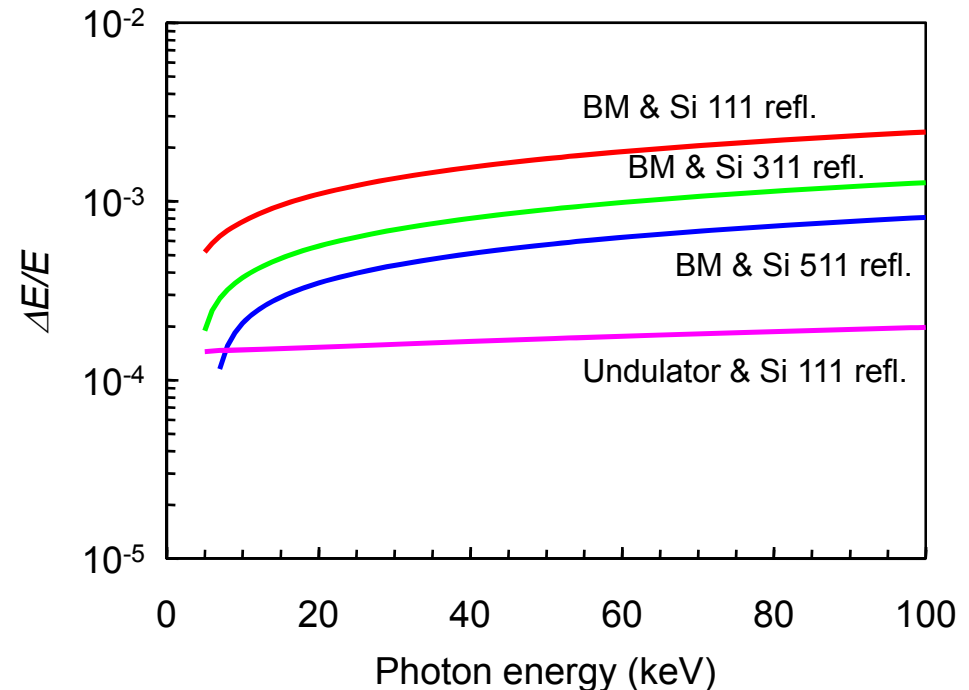
# Energy resolution

$$\frac{\Delta E}{E} = \cot \theta_B \sqrt{\Omega^2 + \omega^2}$$

$\Omega$ : source divergence,  
 $\omega$ : diffraction width

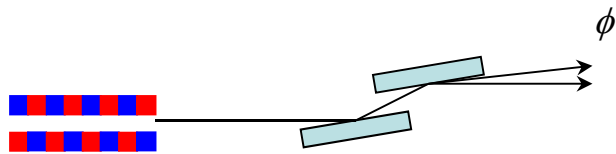


Angle-energy diagram  
 (DuMond diagram)



For usual beamline :  $\Delta E/E = 10^{-5} \sim 10^{-3}$

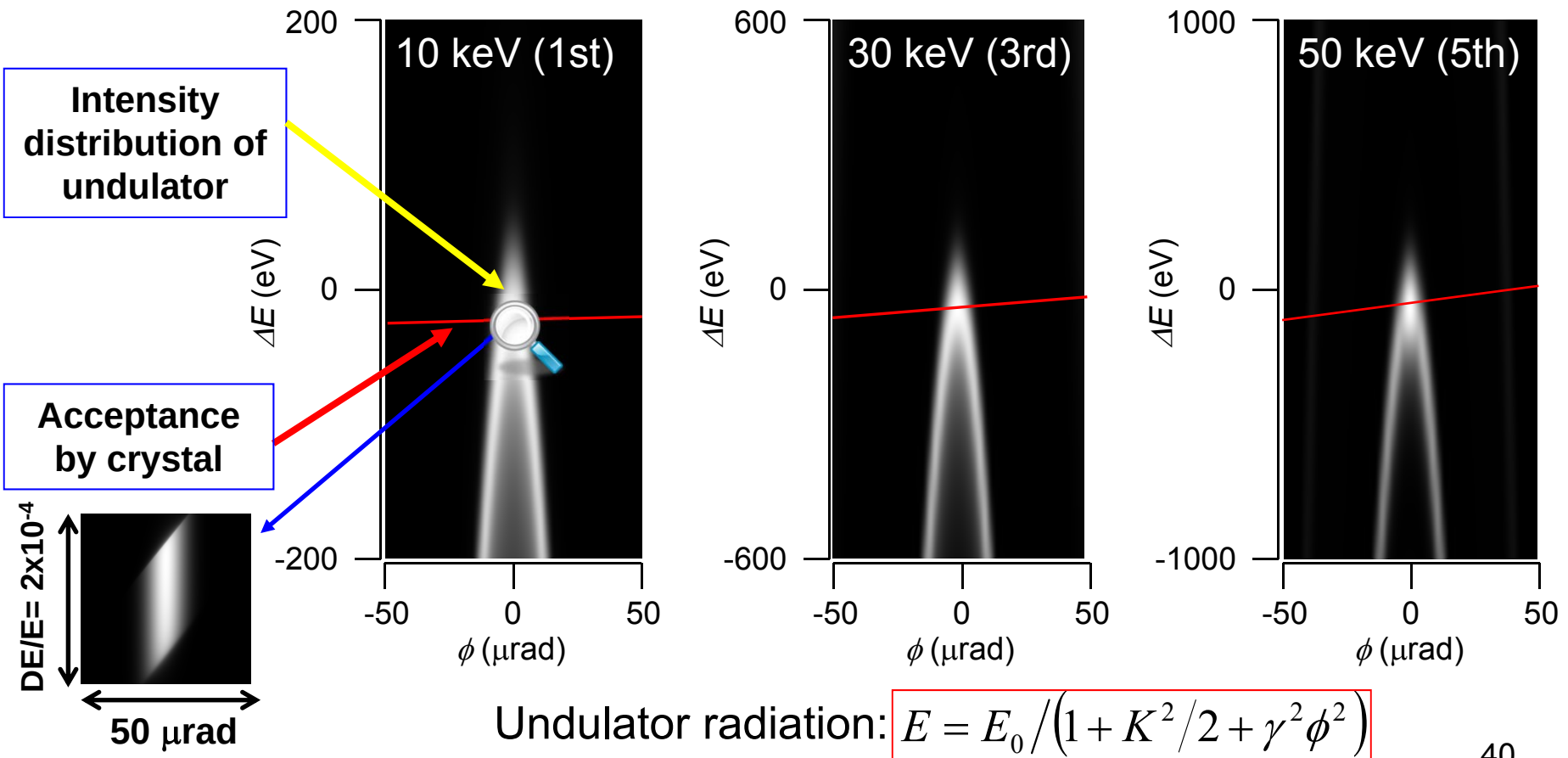
# DuMond diagram: undulator & DCM



SPring-8 standard undulator

( $\lambda_u = 32$  mm,  $N = 140$ ,  $K = 1.34$ ,  $E_{1st} = 10$  keV)

+ DCM (Si 111 refl.)

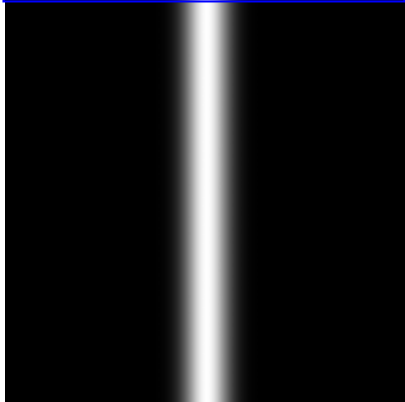


**Wider slit increases unused photons (power) on the monochromator !**

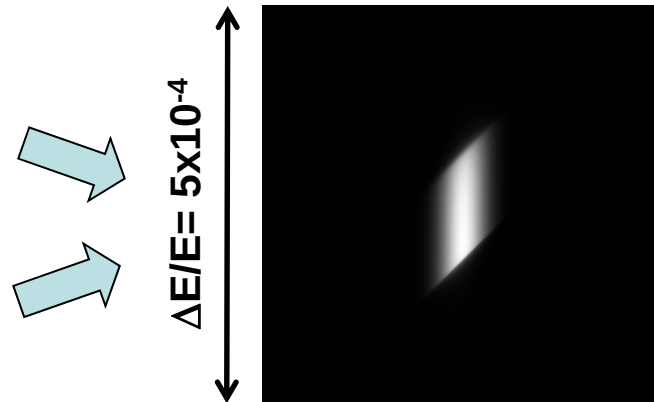
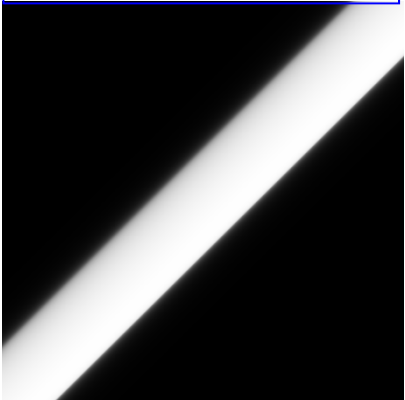


# DuMond diagram: undulator & DCM

Undulator radiation

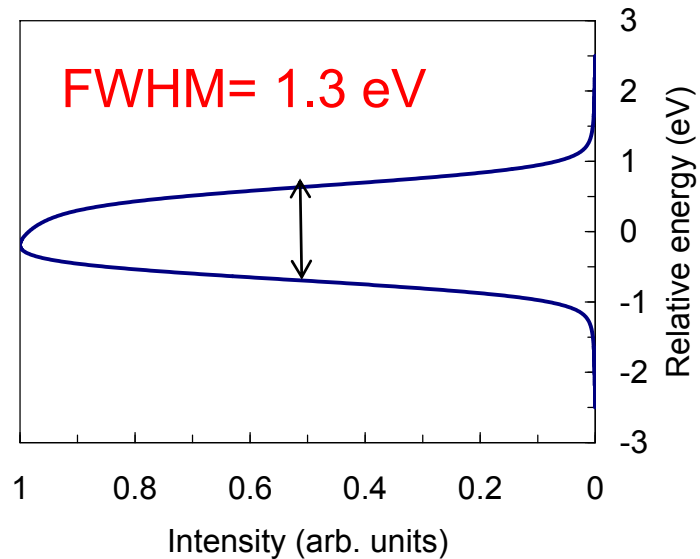


Acceptance by  
Si 111 DCM



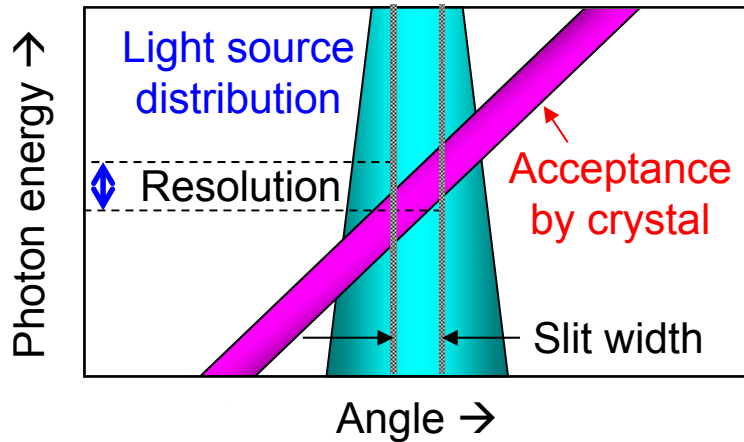
$\Delta E/E = 5 \times 10^{-4}$

100  $\mu$ rad

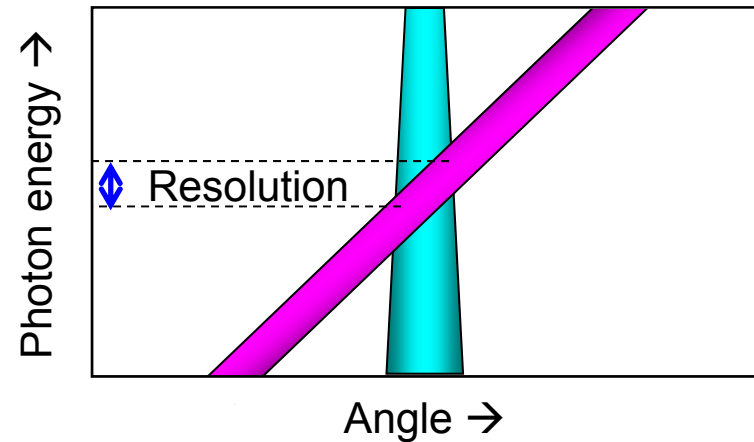


SPring-8 standard undulator + 20  $\mu$ rad slit + Si 111 DCM  
10-keV photons  $\rightarrow 1.3 \times 10^{-4}$

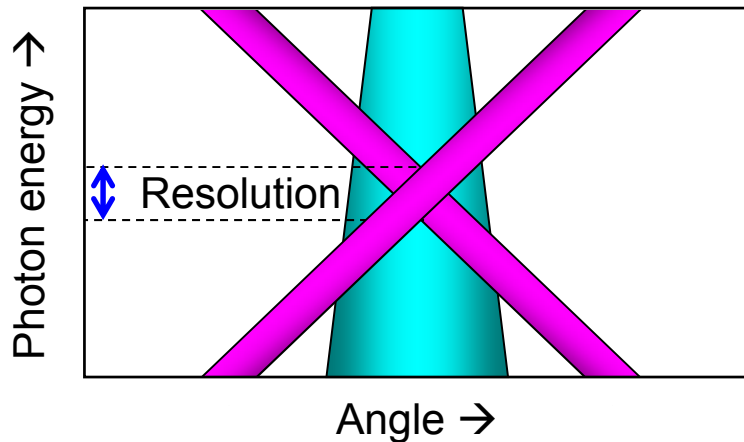
# Improvement of energy resolution



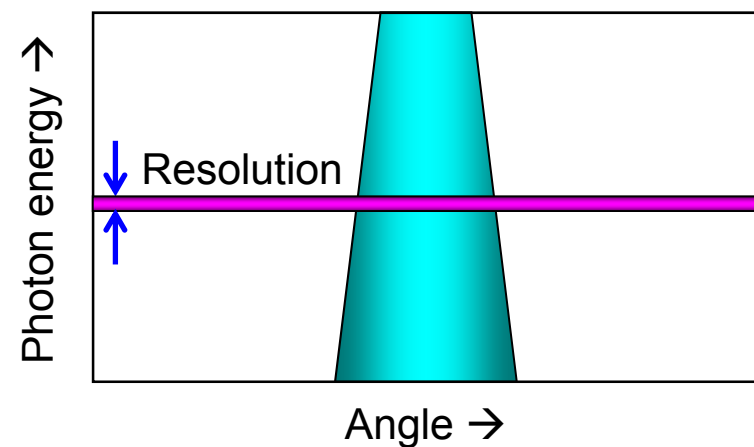
(A) Collimation using slit



(B) Collimation using pre-optics  
w/ collimation mirror, CRL, ..



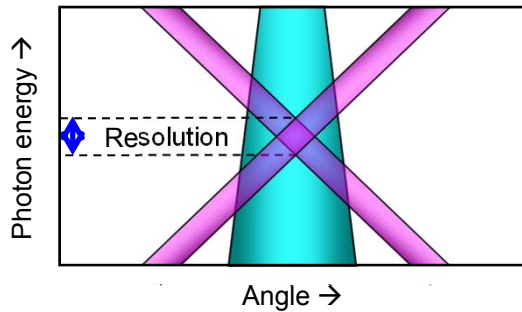
(C) Additional crystal  
w/ (+,+) setting



(D) HR monochromator of  
 $\pi/2$  reflection (~meV)

(B)~(D): restriction on photon energy

# Improvement of energy resolution

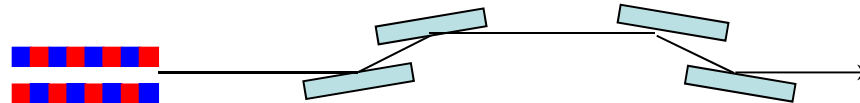


(C) Additional crystal w/ (+,+) setting

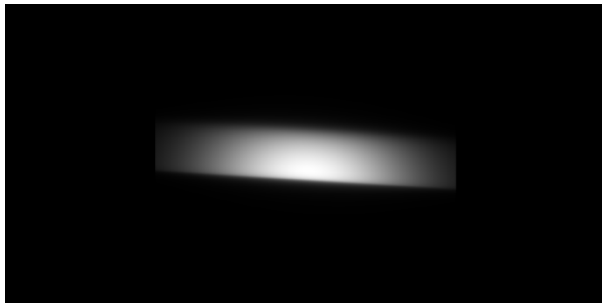
→ HXPES

Si 111 DCM

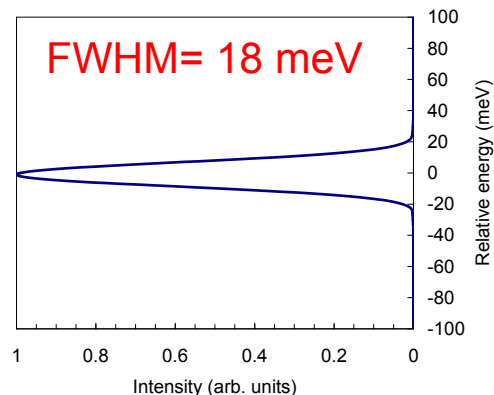
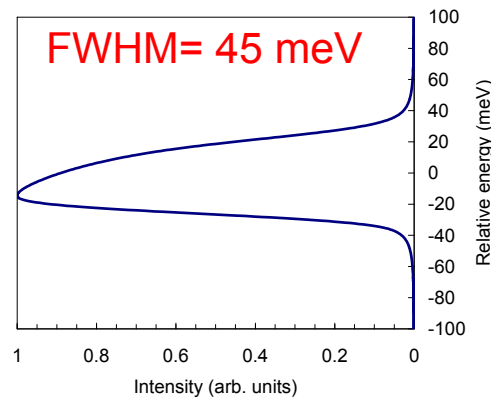
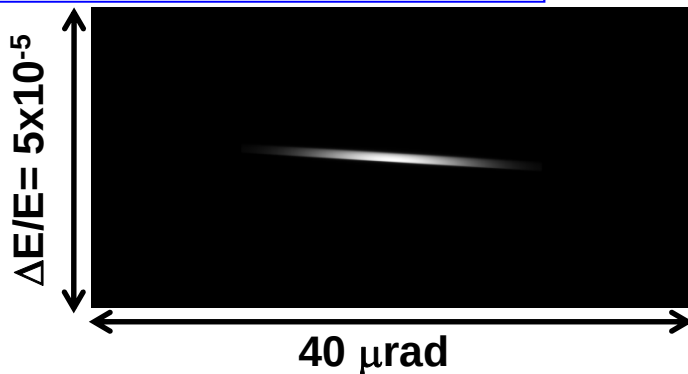
Si *nnn* channel-cut mono.



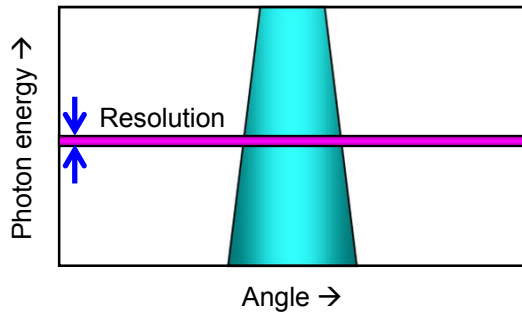
Si 333 refl. for 6 keV



Si 555 refl. for 10 keV

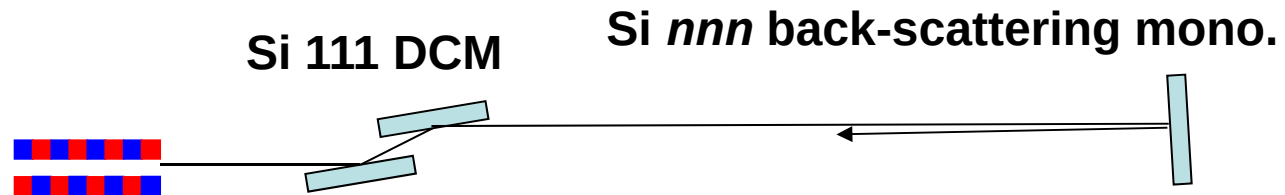


# Improvement of energy resolution

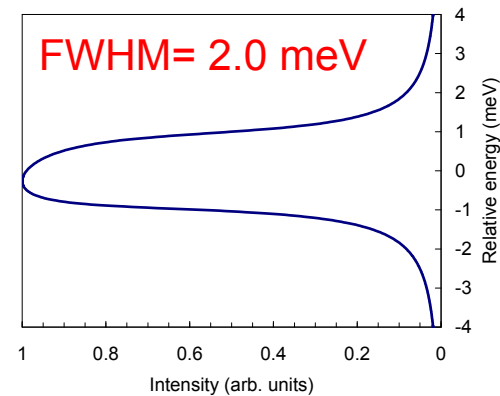
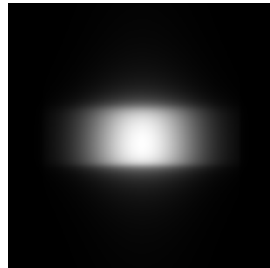


(D) HR monochromator of  $\sim\pi/2$  reflection ( $\sim\text{meV}$ )

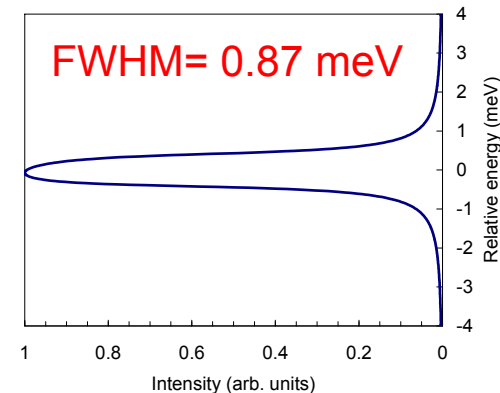
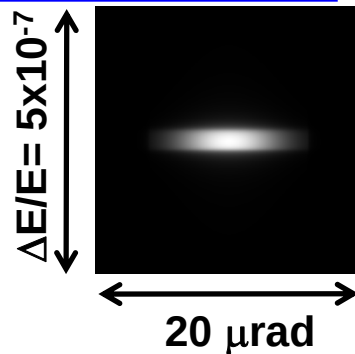
$\rightarrow$  Inelastic scattering



Si 999 refl. for 17.8 keV



Si 11 11 11 refl. for 21.7 keV



# Photon flux after monochromator

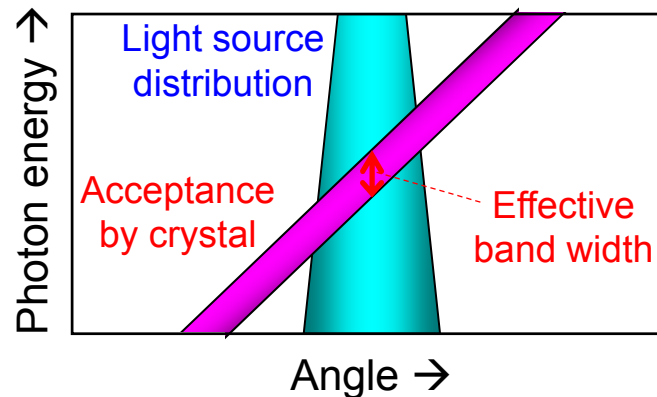
Photon flux (throughput) after monochromator can be estimated using effective band width:

Photon flux (ph/s) =

Photon flux from light source (ph/s/0.1%bw)

x 1000

x Effective band width of monochromator



Throughput is estimated by overlapped area.

Note difference from energy resolution.

# Effective band width

Starting with Darwin width in the energy axis

$$\frac{\Delta E}{E} \approx \frac{|\chi_{hr}|}{\sin^2 \theta_B}$$

$$\chi_{hr} \propto \lambda^2 \{f^0(d_{hkl}) + f'(\lambda)\}$$

Neglecting anomalous scattering factor  $f'$

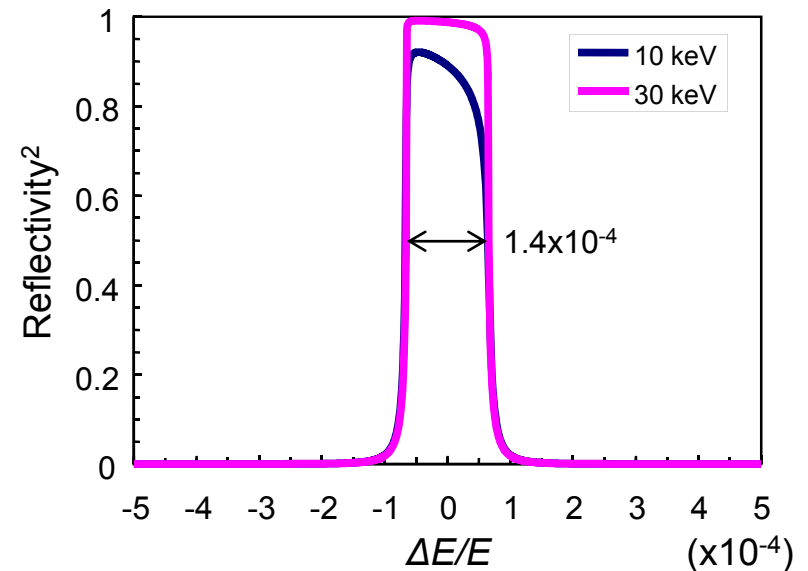
$$\chi_{hr} \propto \lambda^2 f^0(d_{hkl})$$

$$\begin{aligned} \frac{\Delta E}{E} &= -\frac{\Delta \lambda}{\lambda} \approx \frac{|\chi_{hr}|}{\sin^2 \theta_B} \\ &= 4d_{hkl}^2 \frac{|\chi_{hr}|}{\lambda^2} \end{aligned}$$

$$\frac{\Delta E}{E} = -\frac{\Delta \lambda}{\lambda} \propto d_{hkl}^2 f^0(d_{hkl})$$



Independent of photon energy



e.g. for Si 111 refl. DCM case

Note relative energy width is constant.

# Effective band width (Integrated intensity)

For single-bounce monochromator

$$\frac{\Delta E}{E} = \frac{|\chi_{hr}|}{2 \sin^2 \theta_B} \int R(W) dW$$

$$= \frac{8}{3} \cdot \frac{|\chi_{hr}|}{2 \sin^2 \theta_B}$$

↑ For no absorption

For double-crystal monochromator

$$\frac{\Delta E}{E} = \frac{|\chi_{hr}|}{2 \sin^2 \theta_B} \int R(W)^2 dW$$

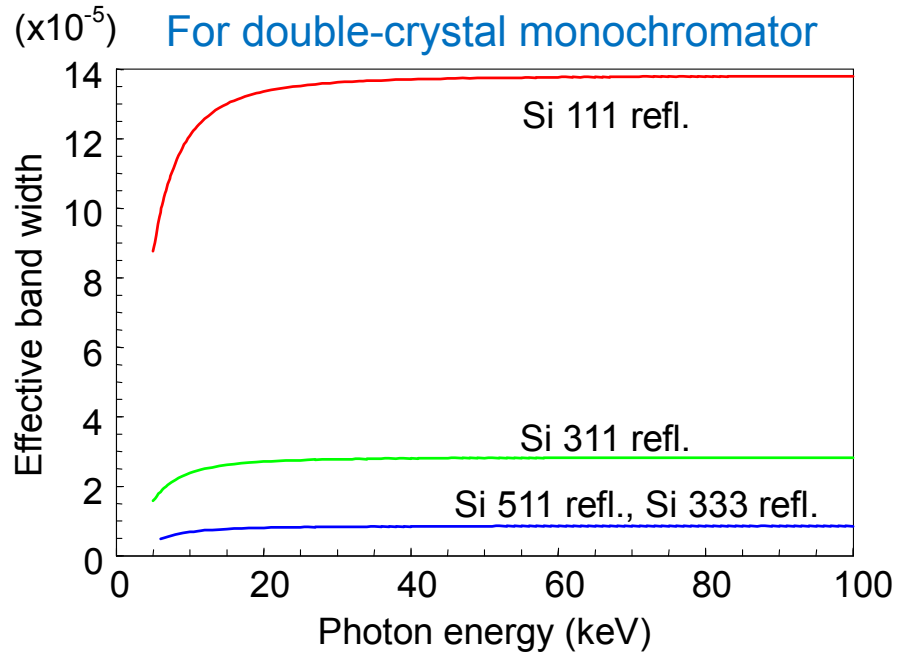
$$= \frac{32}{15} \cdot \frac{|\chi_{hr}|}{2 \sin^2 \theta_B}$$

↑ For no absorption

↑ = ~2

When you need flux → Lower order (Si 111 refl.,...)

When you need resolution → Higher order (Si 311, Si 511 refl.,...)



Effective band-width is obtained by integration of reflection curve.

# Photon flux estimation

## Effective band width

| Reflection (nominal energy) | Effective band width    |
|-----------------------------|-------------------------|
| Si 111 DCM (6 keV)          | $1.0045 \times 10^{-4}$ |
| Si 111 DCM (8 keV)          | $1.1399 \times 10^{-4}$ |

Photon flux (throughput) after monochromator can be estimated using effective band width:

**Photon flux (ph/s) =**

**Photon flux from light source (ph/s/0.1%bw)**

**x 1000**

**x Effective band width of monochromator**

**This approach is valid.**

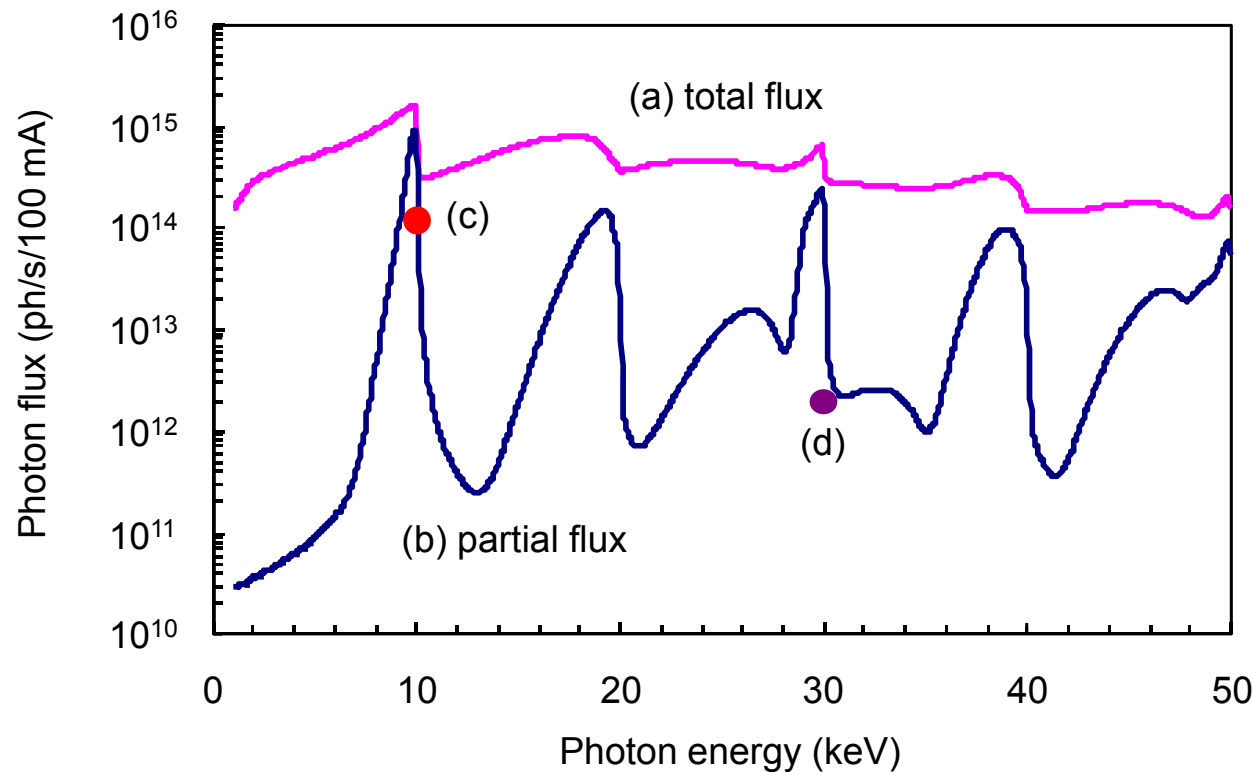
|                     |                       |                       |
|---------------------|-----------------------|-----------------------|
| Si 111 DCM (12 keV) | $5.28 \times 10^{13}$ | $5.29 \times 10^{13}$ |
| Si 111 DCM (14 keV) | $4.20 \times 10^{13}$ | $4.20 \times 10^{13}$ |
| Si 333 DCM (14 keV) | $2.62 \times 10^{12}$ | $2.61 \times 10^{12}$ |

$$(\phi)^2 dEd\phi$$

uMond



# Photon flux at undulator beamline



- (a) Total flux @ 0.1% b.w.
- (b) After frontend slit  
 $1 \times 1 \text{ mm}^2$  @30 m
- (c) Si 111 refl. @10 keV  
Effective b.w. =  $1.3 \times 10^{-4}$
- (d) 3<sup>rd</sup> harmonics @30 keV  
Effective b.w. =  $8.0 \times 10^{-6}$

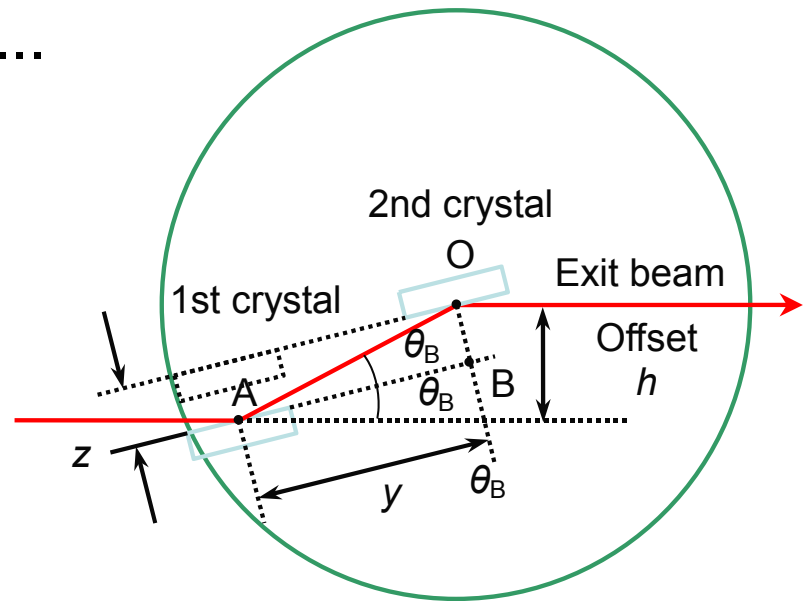
Higher harmonics elimination more → mirror or detuning of DCM

We can obtain photon flux of  $10^{13} \sim 10^{14}$  ph/s/100 mA/mm<sup>2</sup> using standard undulator sources and Si 111 reflections at SPring-8 beamlines.

# Double-crystal monochromator

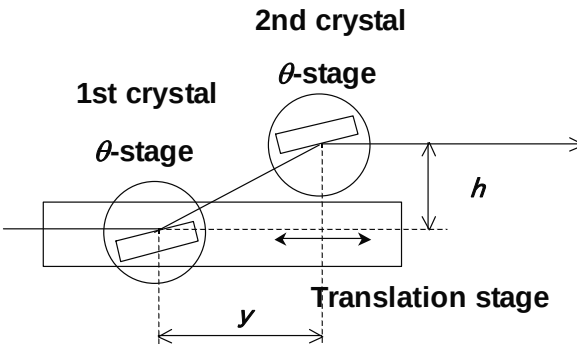
- Fixed-exit operation for usability at experimental station.
- Choose suitable mechanism for energy range (Bragg angle range).
- Precision, stability, rigidity,...

$$y = AB = \frac{h}{2 \sin \theta_B}$$
$$z = OB = \frac{h}{2 \cos \theta_B}$$



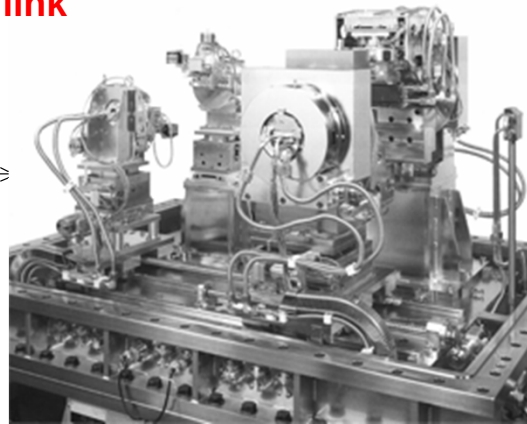
# e.g. Double-crystal monochromator

**$\theta_1$  + translation +  $\theta_2$  computer link**



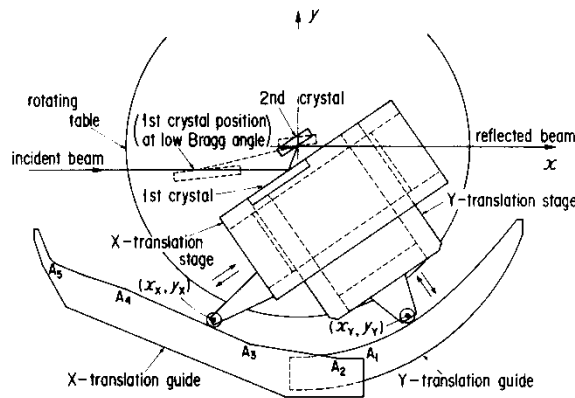
$h = 100 \text{ mm}$ ,

$\theta_B = 5.7 \sim 72^\circ$  (for lower energy range)



SPring-8 BL15XU

**$\theta$  + two translation (2 cams)**



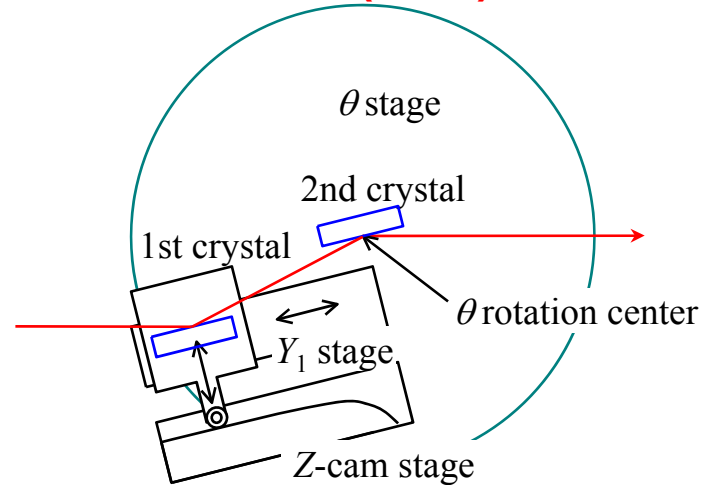
KEK-PF BL-4C

Matsushita et al., NIM A246 (1986)



$h = 25 \text{ mm}$ ,  $\theta_B = 5 \sim 70^\circ$

**$\theta$  + two translation (1 cam)**



Offset  $h = 30 \text{ mm}$   
 $= 3 \sim 27^\circ$  for higher energy range  $\theta_B$

SPring-8 std. DCM

51

Yabashi et al., Proc. SPIE 3773, 2 (1999)

# Crystal cooling

## Why crystal cooling ?

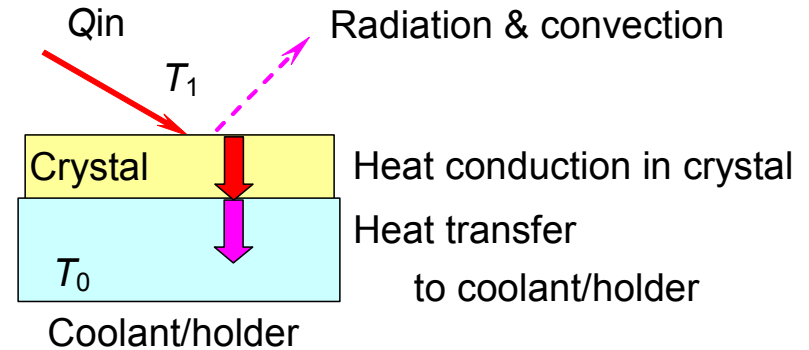
$Q_{in}$  (Heat load by SR) =  $Q_{out}$  (Cooling + Radiation,...)

→ with temperature rise  $\Delta T$

→  $\alpha \Delta T = \Delta d$  ( $d$ -spacing change)

$\alpha$ : thermal expansion coefficient

or →  $\Delta \theta$  (bump of lattice due to heat load)



## Miss-matching between 1st and 2nd crystals occurs:

→ Thermal drift, loss of intensity, broadening of beam, loss of brightness

→ Melting or limit of thermal strain → **Broken !**

### We must consider:

- Thermal expansion of crystal:  $\alpha$ ,
- Thermal conductivity in crystal:  $\kappa$ ,
- Heat transfer to coolant and crystal holder.

### Solutions:

(S-1)  $\kappa/\alpha$  → Larger

(S-2) Contact area between crystal and coolant/holder  
→ larger

(S-3) Irradiation area → Larger,  
and power density → smaller

## Figure of merit

|                             | Silicon              | Silicon             | Diamond            |
|-----------------------------|----------------------|---------------------|--------------------|
|                             | 300 K                | 80 K                | 300 K              |
| $\kappa$ (W/m/K)            | 150                  | 1000                | 2000               |
| $\alpha$ (1/K)              | $2.5 \times 10^{-6}$ | $-5 \times 10^{-7}$ | $1 \times 10^{-6}$ |
| $\kappa/\alpha \times 10^6$ | 60                   | 2000                | 2000               |

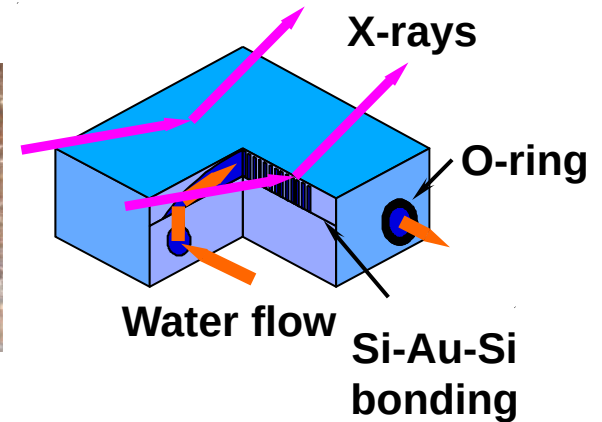
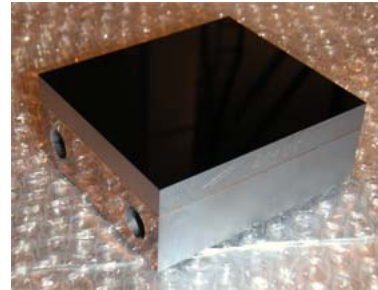
Figure of merit of cooling:  
Good for silicon (80 k)  
and diamond (300 K)

# Crystal cooling at SPring-8

## <Bending magnet beamline>

Power & power density:  
~100 W, ~1 W/mm<sup>2</sup>

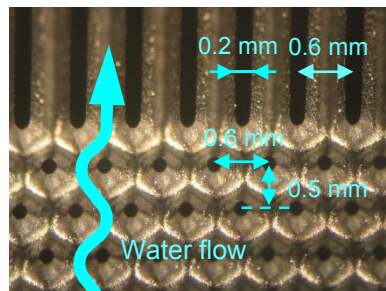
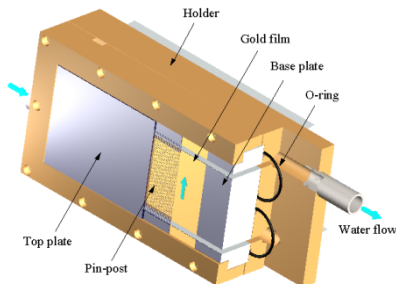
### Fin crystal direct-cooling - (S2)



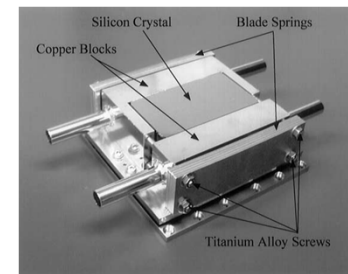
## <Undulator beamline>

Linear undulator,  $N=140$ ,  $\lambda_u=32$  mm  
Power & power density: 300~500 W ,  
300~500 W/mm<sup>2</sup>

### a) Direct cooling of silicon pin-post crystal – (S2) & (S3)



### b) Silicon cryogenic cooling - (S1)



### c) Ila diamond with indirect water cooling - (S1)



# Summary

---

Key issues on the x-ray **monochromator** were shown,  
introducing the **dynamical x-ray diffraction** for **large & perfect crystal**,  
w/ several important points:

- 1) Total reflection occurs at the gap between dispersion surfaces.
- 2) Normalized deviation parameter  $W$  is related to the gap.
- 3)  $W$  is parameter of angular deviation and energy (wavelength) deviation.  
It gives **DuMond diagram** as a band of  $|W| < 1$ .
- 4) By combination of light source and monochromator crystals,  
photon energy, energy resolution, photon flux, · · · can be controlled / tuned.

Double-crystal monochromator w/ crystal cooling is needed for practical use  
at the SR beamline.

By understanding these, you will be approaching to good design/use of the beamline  
for your SR science.



# Text books following Laue's dynamical theory

Ergebnisse der Exakt Naturwiss.

10 (1931) 133-158.

## Die dynamische Theorie der Röntgenstrahlinterferenzen in neuer Form.

Von M. v. LAUE, Berlin.

Mit 4 Abbildungen.

Inhaltsverzeichnis.

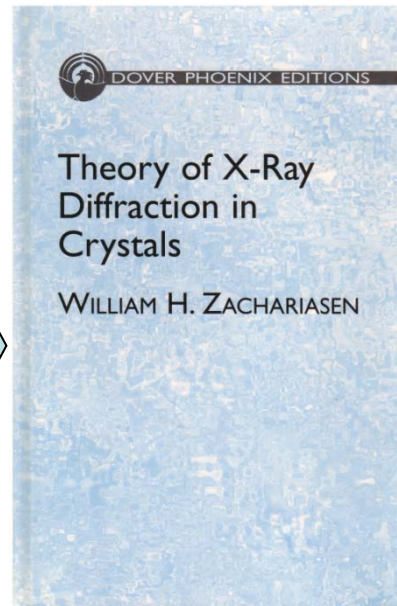
|                                                                                              |           |
|----------------------------------------------------------------------------------------------|-----------|
| § 1. Die dynamische Grundgleichung . . . . .                                                 | Seite 137 |
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EWALD'S dynamische Theorie der Röntgenstrahlinterferenzen<sup>1</sup> gehört nach unserer Ansicht auf alle Zeiten zu den Meisterwerken der mathematischen Physik. Sie bewältigte mit glänzenden Methoden ein zunächst schier unlösbares Problem, rechtfertigte eigentlich erst die elementare, rein auf Phasensummensetzung beruhende Theorie. Die Unzulänglichkeiten quantitativ abzuschätzen lehrte, die Verbundenheit mit der optischen Dispersionstheorie, die Abweichungen von den Ergebnissen der älteren Theorie. Wenige Jahre später für die vollständige Deutung des HJALMAR<sup>2</sup> sowie für eine exakte Messung der Wellenlänge wurden. Aber wir glauben aussprechen zu dürfen, dass die ihre Resultate oft benutzen, nicht allzu durchgearbeitet haben. Denn jene glänzenden mathematischen Methoden waren nicht nur schwierig zu finden, sie bereiten in manchen Teilen auch dem Leser Schwierigkeiten. Duran hat auch die Neubearbeitung durch WALLER<sup>3</sup> nicht viel geändert, welche im übrigen das Verdienst besitzt, zuerst Gitter mit Basis in die Rechnung einbezogen zu haben. Der Zweck der folgenden Seiten ist, der Theorie eine leichter verständliche Form zu geben. Daß wir trotz weitgehenden Verzichts auf neue Ergebnisse die neue Form veröffentlichten, scheint uns durch die Wichtigkeit der Sache gerechtfertigt; ebenso, daß wir, um dem Leser das Zurückgreifen auf die genannten Arbeiten zu erleichtern, die geometrische Diskussion EWALD'S nochmals zum Abdruck bringen, obwohl wir an ihr fast nichts zu ändern brauchen.

<sup>1</sup> EWALD, P. P.: Ann. Physik 54, 519, 557 (1917); Z. Physik 2, 335 (1920); 30, 1 (1924).

<sup>2</sup> WALLER, J.: Uppsala Universitets Årsskrift 1925.

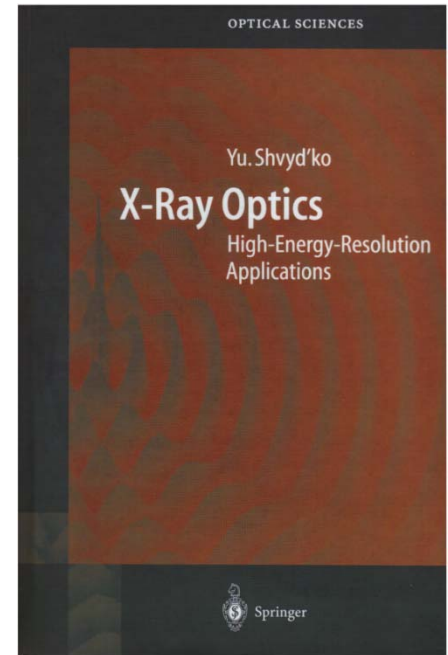
Dover (1945)



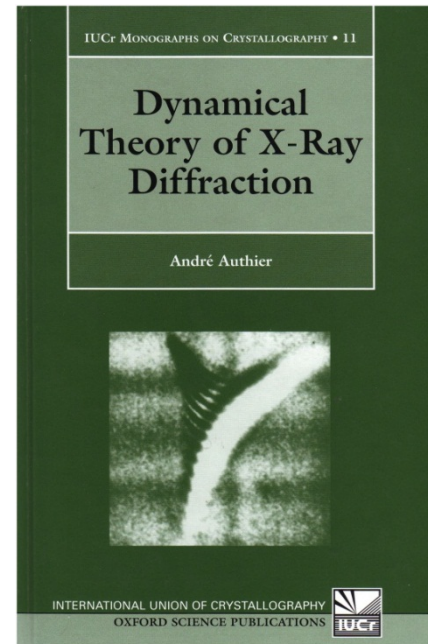
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R. W. James

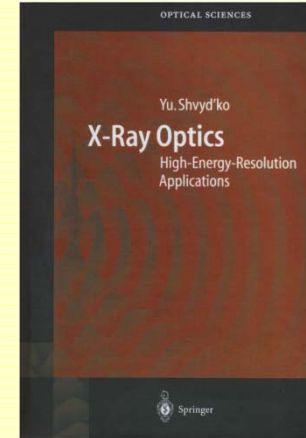
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***Thank you for your attention.***